Atlas of Procedures in Neonatology









Mhairi G. MacDonald Jayashree Ramasethu



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Editors: MacDonald, Mhairi G.; Ramasethu, Jayashree Title: Atlas of Procedures in Neonatology, 4th Edition Copyright ©2007 Lippincott Williams & Wilkins > Front of Book > Dedication Dedication "This book is dedicated to my mother, Jane Graham MacDonald, for her steadfast support, love, and encouragement.â€ Mhairi G. MacDonald MBChB,FRCPE,FRCPCH,DCH

Editors: MacDonald, Mhairi G.; Ramasethu, Jayashree Title: Atlas of Procedures in Neonatology, 4th Edition Copyright ©2007 Lippincott Williams & Wilkins > Front of Book > Preface Preface

It has been over a quarter of a century since the first edition of the Atlas of Procedures in Neonatology was published. The basic philosophy and purpose of the Atlas, as reflected in the preface to editions one through three, has not changed as the field of neonatal-perinatal medicine has matured.

This fourth edition has undergone a significant facelift, and the majority of illustrations are now in color. Color photographs of procedure complications proved comparatively difficult to obtain; it is tempting to hope that this reflects a decline in their incidence. The new, fold-over cover continues to allow the book to open flat and also offers easy identification on a bookshelf.

In the preface of the third edition, we noted that some promising technologies, such as transcutaneous bilirubin measurement, required further field testing. A chapter on transcutaneous bilirubin measurement has been added in this edition, plus one on auditory screening, and three other new chapters.

With the exception of Aseptic Preparation, the procedures included on the DVD fall into two categories:

• Commonly performed procedures, such as peripheral and umbilical line placement and endotracheal intubation.

• Vital emergency procedures that trainees may have infrequent opportunity to perform, such as chest aspiration/placement of a thoracostomy tube, or rare opportunity to perform, such as exchange transfusion.

Particularly in the case of exchange transfusion, there is now a generation of neonatologists who have graduated from training with limited practical experience of the procedure but who now have the responsibility to teach it to trainees. We hope that the videoâ€"animation will prove a valuable teaching resource.

Mhairi G. MacDonald MBChB, FRCPE, FRCPCH, DCH Jayashree Ramasethu MD, FAAP

Editors: MacDonald, Mhairi G.; Ramasethu, Jayashree Title: Atlas of Procedures in Neonatology, 4th Edition Copyright ©2007 Lippincott Williams & Wilkins > Front of Book > Preface to the Third Edition Preface to the Third Edition

Almost two decades have passed since the publication of the first edition of Atlas of Procedures in Neonatology. We have seen our patients become progressively smaller, less mature, and more susceptible to iatrogenic morbidity. The increasing fragility of our patient population and the escalating complexity of care continue to make neonatology challenging. These challenges have necessitated not only innovations in technology and equipment but also increased familiarity with and proficiency in procedures.

This revised edition of Atlas of Procedures in Neonatology reflects changes that have come about in the management of neonates since the second edition was published in 1993. Reliable monitoring techniques and therapeutic modalities have been developed or refined since then for some aspects of care; however, other promising technologies, such as noninvasive transcutaneous bilirubin monitoring, require additional field testing and validation.

The passing of the era of the mercury thermometer deserves special note. The prototype of the widely used clinical mercury thermometer was designed over 130 years ago, an admirable track record for a medical device. Technical advances in thermometry have rendered the risk of mercury toxicity from broken thermometers unacceptable. As a result, in July 2001, the Committee on Environmental Health of the American Academy of Pediatrics recommended that mercury thermometers no longer be used for pediatric patients.

Advances in the prevention and intrauterine management of alloimmune hemolytic disease, along with the liberal use of phototherapy, have resulted in a significant decrease in the need for postnatal exchange transfusionsâ€"to the point that fellows in neonatalâ€"perinatal medicine now occasionally complete training programs without ever performing the procedure. The reappearance of kernicterus in well full-term neonates underscores the importance of retaining the chapter on exchange transfusions in this edition. Cryotherapy for retinopathy of prematurity has been replaced by laser therapy. A chapter on ostomy care has been added. Expanded chapters on the management of extravasation injuries and vascular spasm and thrombosis are acknowledgments of the all-too-common iatrogenic problems in neonatal intensive care. A new chapter on perimortem sampling provides guidelines for testing in the event of sudden or unexpected death.

Updated information regarding the complications of each procedure is provided to facilitate risk-versusbenefit considerations and the informed consent process. Complications are listed whenever possible in order of frequency of occurrence or of importance.

At the request of those who used the previous editions of Atlas of Procedures in Neonatology, we have modified the binding to allow the book to lie flat when open. As in previous editions, commercial products listed in the text are intended for illustrative purposes only; no endorsement is implied. Commercial availability of some equipment may change subsequent to the publication of this edition.

Acknowledgment of the fact that invasive procedures are currently unavoidable in the care of sick neonates must come with the recognition that there is responsibility to reduce potential iatrogenic morbidity to the

minimum possible. The latter may be achieved by strict adherence to basic principles of asepsis, careful monitoring and maintenance of patient homeostasis, and by ensuring that procedures are performed or supervised by those with the requisite expertise. Mhairi G. MacDonald MBChB, FRCP(E), DCH

Jayashree Ramasethu MD

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There is nothing more gratifying to editors than to see their book lying open and dog-eared during a bedside visit to another facility. We have been pleased to find the first edition of the Atlas of Procedures in Neonatology used not only by trainees and staff in neonatology but also by other members of the neonatal–perinatal health care team, including radiologists and respiratory therapists. The primary purpose of the Atlas is to provide a detailed, step-by-step approach to procedures, most of which are performed by neonatologists, pediatricians, and nurses within the nursery. Some proceduresâ€"such as extracorporeal membrane oxygenation cannulation, operative tracheotomy, gastrostomy, and cryotherapyâ€"are usually performed by surgical specialists but are included to promote understanding by those who are responsible for the perioperative care of the neonate. On the advice of our readers, we have selected a binding that will allow the book to open flat so as to facilitate bedside use during procedures. The organizational format of the first edition remains. We recommend studying an entire procedure before starting it, not only to review the technique but also to better weigh benefits against risks by understanding the complications and precautions. As in the first edition, we have emphasized the anatomical differences between the neonate and older patients that influence the performance of certain procedures. After every procedure, we have attempted to include a comprehensive list of complications in order to heighten awareness of their potential impact on both morbidity and mortality. The order of listing does not necessarily reflect the frequency or severity of any single complication.

It is sobering to observe that a significant number of complications, some of them not previously recognized, continue to be reported for procedures that have been standard in neonatal nurseries for more than two decades. For example, since the first edition of the Atlas was published, reports in the literature on complications of umbilical artery catheterization have approximately tripled. With every new procedure there is a learning curve, but one would expect the incidence of complications to decrease as experience and expertise increase. Clearly, the number of reported complications does not represent their true incidence. An optimistic view might be that increased reporting of complications reflects a more universal respect for the possibility of their occurrence and attempts to find ways of preventing or minimizing them. When any procedure is applied to smaller and more immature infants, it is not only technically more difficult but also more likely to be accompanied by side effects or complications. For all procedures performed in the newborn there is a baseline morbidity; no procedure will be absent complications. For example, placement of a peripheral intravenous line is a basic procedure essential for the survival of sick

newborn babies. There have been significant improvements in the size and quality of i.v. cannulas and pumps specifically to allow for pressure obstruction alarms, low flow rates, and so forth. However, no matter how good the care of the infant and the i.v., there will always be incidents of infiltration and chemical skin burns. It behooves each clinician to carefully weigh the risks versus the benefits of every procedure before beginning it, while any piece of equipment remains in place, and even in the months and years after completion.

One cannot possibly practice good medicine and not understand the fundamentals underlying therapy. Few if any rules for therapy are more than 90% correct. If one does not understand the fundamentals, one does

more harm in the 10% of instances to which the rules do not apply than one does good in the 90% to which they do apply. â€"Fuller Albright Mary Ann Fletcher MD Mhairi G. MacDonald MBChB, FRCP(E), DCH

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The rapid advances in neonatology in the last 15 years have brought with them a welter of special procedures. The tiny, premature, and the critically ill term neonate is attached to a tangle of intravenous lines, tubes, and monitoring leads. As a result, more and more procedures are done at the bedside in the intensive-care nursery, rather than in a procedure room or operating room. With these technical advances has come the opportunity for more vigorous physiologic support and monitoring. With them also has come a whole new gamut of side-effects and complications. The old dictum to leave the fragile premature undisturbed is largely ignored. It is therefore the responsibility of those who care for sick newborns to understand the complications as well as the benefits of new procedures and to make systematic observations of their impact on both morbidity and mortality. Unfortunately, the literature on outcome and complications of procedures is widely scattered and difficult to access. Manuals that give directions for neonatal procedures are generally deficient in illustrations giving anatomic detail and are often cursory. We are offering Atlas of Procedures in Neonatology to meet some of these needs. A step-by-step, practical approach is taken, with telegraphic prose and outline form. Drawings and photographs are used to illustrate anatomic landmarks and details of the procedures. In several instances, more than one alternative procedure is presented. Discussion of controversial points is included, and copious literature citations are provided to lead the interested reader to source material. A uniform order of presentation has been adhered to wherever appropriate. Thus, most chapters include indications, contraindications, precautions, equipment, technique, and complications, in that order.

The scope of procedures covered includes nearly all those that can be performed at the bedside in an intensive-care nursery. Some are within the traditional province of the neonatologist or even the pediatric house officer. Others, such as gastrostomy and tracheostomy, require skills of a qualified surgeon. Responsibility for procedures such as placement of chest tubes and performance of vascular cutdowns will vary from nursery to nursery. However, some details of surgical technique are supplied for even the most invasive procedures to promote their understanding by those who are responsible for sick neonates. We hope this will help neonatologists to be more knowledgeable partners in caring for babies and will not be interpreted as a license to perform procedures by those who are not adequately qualified.

The book is organized into major parts (e.g., $\hat{a} \in \mathbb{C}$ Vascular Access, $\hat{a} \in \mathbb{C}$ Tube Placement, $\hat{a} \in \hat{a} \in \mathbb{C}$ Respiratory Care $\hat{a} \in \mathbb{C}$), each of which contains several chapters. Most chapters are relatively self-contained and can be referred to when approaching a particular task. However, Part I, $\hat{a} \in \mathbb{C}$ Preparation and Support, $\hat{a} \in \mathbb{C}$ is basic to all procedures. Occasional cross referencing has been used to avoid repetitions of the same text material. References appear at the end of each part.

Many persons have contributed to the preparation of this atlas, and we are grateful to them all. Some are listed under Acknowledgments, and others have contributed anonymously out of their generosity and good will. Special thanks is due to Bill Burgower, who first thought of making such an atlas and who has been gracious in his support throughout this project.

If this atlas proves useful to some who care for sick newborns, our efforts will have been well repaid. Neonatology is a taxing field: strenuous, demanding, confusing, heartbreaking, rewarding, stimulating, scientific, personal, philosophical, cooperative, logical, illogical, and always changing. The procedures described in this atlas will eventually be replaced by others, hopefully more effective and less noxious. In the meantime, perhaps the care of some babies will be assisted. Mary Ann Fletcher MD Mhairi G. MacDonald MBChB, FRCP(E), DCH Gordon B. Avery MD, PhD

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1-Informed Consent for Procedures on Neonates

Harold M. Ginzburg

A. General Comments

Medicine is an art and a science needed by most, feared by some, and observed by all. Modern medical care in hospitals is generally provided by "teams†of medical personnel including but not limited to physicians, nurses, therapists, and technicians (e.g., extracorporeal membrane oxygenation, respiratory therapy, dialysis) (1,2 and 3). By medical tradition, the physician and the hospital have been held responsible for the actions of the team members. Today, each team member is also medically and legally responsible for his or her own actions.

It is the fundamental duty and responsibility of those who provide medical care and treatment to neonates to inform the parent or guardian appropriately and to document that relevant information has been provided, and understood, and that informed consent has been obtained. Failure to obtain fully informed consent may result in legal liability for the health care provider under either a claim of negligence or a claim of assault and battery (depending on state law) (4,5,6,7).

Neonatologists, obstetricians, and pediatricians practice in a particularly emotionally charged area of medicine. Families are more ready to accept mortality and morbidity in their senior members than in their youngest members. Options for treatment must be carefully explained and include the option, when appropriate, of no treatment. Options for treatment should be presented in terms of the immediate, intermediate, and long-term effects of the treatment on the neonate and on those who will be directly affected by the medical decisions.

The federal government and the individual states and territories have laws, regulations, guidelines, policies, and practices that directly and indirectly affect the practice of medicine care (8,9). Federal and state reimbursement, licensure, inspection, and enforcement functions vary and may even be in conflict with each other. Therefore, health care providers are encouraged to consult with hospital administrative personnel and legal counsel when potential legal issues arise because, in the final analysis, $\hat{a} \in \alpha$ The law does not permit [a health care provider] to substitute his own judgment for that of the patient [or guardian] (10). $\hat{a} \in \alpha$

B. Medical Legal Concepts

- Duty
 - Legal: A duty is a legal and ethical responsibility. A breach (violation) of a duty owed to another person may result in legal liability. Duties are a reflection of the moral fiber of a community; they are codified in state and federal statutes and regulations. Public policies and laws thus become the vehicle for expressing a community's moral imperatives.
 - Medical: A health care provider has a duty to conform his or her practice to a reasonable (or, in some instances, usual) standard of care specific to the type of illness and to the particular circumstances in which the care and/or treatment are being provided. This duty includes performing only procedures for which, under normal circumstances, appropriate equipment and support are available and that he or she possesses the competence to perform.
- Health care provider–patient relationship
 - Fiduciary ("pertaining to or involving one who holds something in trust for anotherâ€): Health care providers and hospital care systems have a fiduciary duty to patients and their parents or guardians. A fiduciary duty is a responsibility that arises from the trust and confidence placed by the patient/parent/guardian in the health care professional. Health care providers are assumed to have superior medical knowledge; they are expected to warn their patients and to attempt to protect them from untoward or predictable harm. The fiduciary relationship is created when a health care provider responds to an expressed or implied request for treatment by the patient, his or her guardian, or a third party (e.g., emergency medical service personnel); the health care provider then has a duty to share his or her

knowledge about the nature of the illness, its prognosis, treatment options, and associated risks (these are the essential elements of an informed consent agreement). The health care provider also has a duty to protect patient information from access by those who do not have a legal right to access it.

- Contract: The health care provider–patient relationship is a contract. A health care provider and a patient (or his or her parent or guardian) enter into the contract, either implied or expressed, either verbally or written, for the performance of medical services. The contract created is based on the fiduciary relationship and not on a financial one.
- Standard of care
 - Legal: The health care provider has a duty to conform his or her practice to a reasonable standard of care in the particular type of case and in the particular circumstances in which the care and treatment are being provided.
 - Medical: The health care provider must not undertake any procedure that will place the patient at an unreasonably great risk of harm when weighed against the potential benefits. The health care provider must consider the consequences of his or her actions, or inactions, and exercise his or her best judgment when providing care.
- Negligence
 - General negligence: Negligence is the failure to do something that a reasonable man or woman, guided by those considerations that ordinarily regulate the conduct of human affairs, would do or the doing of something that a prudent and reasonable man or woman would not do.
 - Medical negligence: Medical negligence, or medical malpractice, is a special instance of negligence. A missed operative diagnosis, a therapeutic misadventure, a failure to arrange for follow-up care, and a failure to inform, warn, or protect the patient, or in some instances a third party, may be special instances of medical negligence. The medical care profession is held to a specific minimum level of performance based on the possession, or claim of possession, of "special knowledge or skills†that have accrued through specialized education and training. A health care provider may be sued successfully only if there was a violation of a duty or obligation, recognized by law, that required the defendant-health care provider to conform to a particular standard of conduct in order to protect others, usually the patient, against unreasonable risks of harm.
- Informed consent
 - Clinical: Informed consent is educated consent; it is a cornerstone in the provision of medical care (11). Fundamentally, informed consent is a contract between the physician and the patient and/or the parent or guardian. Informed consent requires that sound, reasonable, comprehensible, and relevant information be provided by a health care professional to a competent individual (patient and/or their parent(s) or guardian) for the purpose of eliciting a voluntary and educated decision about the advisability of permitting one course of clinical action as opposed to another (12). Thus, informed consent involves (a) a clearly stated, understandable offer by the health care provider to provide services and (b) acceptance of the services by the patient or the surrogate; it is also (a) an offer, by the patient or the surrogate, to pay for the services or at least to acknowledge that they have a worth, and (b) it is the health care provider's acceptance of the payment or acknowledgment (13). Informed consent should also prevent unrealistic expectations from evolving, because all parties are made aware of the possibility of the failure of the proposed treatment plan.

Informed consent implies that not only the treating physician, but all those who are involved in the patient's care (e.g., nurse practitioners, nurses, respiratory therapists) have a responsibility to use sound judgment and provide quality care (14). To create a valid contract, there is a legal requirement that an individual providing consent be legally competent and have the legal capacity or ability to enter into that contract (15). Health care providers treating children are sometimes faced with emotionally immature, incoherent, uncooperative, absent, or intoxicated parents. In such instances, assistance from the hospital attorneys and the courts may be required.

Research: The Declaration of Helsinki, agreed to by the World Medical Association in 1964, distinguished between "clinical research combined with professional care†(i.e., research that might directly benefit the patient) and "nontherapeutic clinical research (16).†Special and specific informed consent needs to be obtained for procedures conducted as part of research studies unless the procedures are considered to be a component of the "routine medical care†(see next section). The U.S. Department of Health and Human Services (HHS) (17) and the U.S. Food and Drug Administration (FDA) (18) have promulgated a series of regulations to ensure that research subjects, especially children, are adequately protected when they are enrolled in clinical trials or other clinical experiments. HHS has the power to investigate and sanction investigators for violating its regulations (19).

C. Obtaining Informed Consent

- The person obtaining consent should be aware that there are two levels of informed consent.
 - General informed consent (often referred to as a "blanket consentâ€): The parent or \circ guardian must understand that admission to a hospital entails active clinical intervention by a number of health care providers. While the parent or guardian can expect explanations of many of the procedures that will be used to help his or her child or ward, he or she cannot expect to be informed about every intervention. Routine medical care is considered to be covered under a general informed consent; each time blood is drawn or a nonexperimental medication is administered, a specific informed consent is not required. Each medical facility needs to define (within the limits of what is reasonable for the type of care provided as well as what is reasonable given the social context of the community) the components of routine care. For instance, in an intensive care nursery, placing an arterial line in a critically ill infant for the purpose of obtaining blood gases is not a unique procedure requiring a specific informed consent, whereas placing a neonate on extracorporeal membrane oxygenation would generally require specific informed consent. It is the procedures in the "grayâ€ area that require definition by the facility and individual unit as standard or otherwise. Both a lumbar puncture and the percutaneous placement of a central venous catheter may be considered standard routine procedures in a critical care unit. However, on a general pediatric ward, the percutaneous placement of a central close-up venous line might require specific informed consent.
 - Specific informed consent: The parent or guardian must recognize that beyond the routine medical care provided in a neonatal intensive care unit (NICU), there are specialized medical and surgical procedures that require that they be provided with specific additional information. This information will assist them in determining whether they should consent to the recommended procedure(s) on behalf of their infant. For each such procedure, a written description or an oral description documented in the patient's chart, describing the procedure and the risks and benefits of performing the procedure versus another procedure versus not performing the procedure should be provided.
- The person obtaining the informed consent should understand the following general concepts.
 - Coercion or undue influence: The health care provider has a duty to ensure that the parent or guardian voluntarily assents to permit the treatment or clinical research. Implied threats (such as the unintended impression that the quality of the health care provided by the team to the infant will be less if the procedure is refused), or inducements (no matter how apparently insignificant, e.g., the offer of a transport subsidy contingent on their agreement to the procedure) have the potential to influence the decision and are unacceptable.
 - Mental capacity of the parent or guardian: The parent or guardian is presumed to be competent and have the capacity to understand the medical information, to remember that

information, and to make logical inferences and conclusions from the information. If there is a definite indication that the parent or guardian is not competent, then the health care provider should not accept consent. At that time, the hospital attorney and the local court may become intimately involved in the case.

- The level of understanding of the parent or guardian: The information must be provided in language that the parent or guardian can understand. A qualified medical interpreter should be available if English is not the parent or guardian's first language; use of untrained personnel may lead to miscommunication.
- The risk/benefit for a given therapeutic intervention: The parent or guardian must be provided with information regarding the frequency and severity of the adverse potential consequences as compared to the likelihood, duration, and degree of anticipated benefit from the treatment(s). Where relevant benefits are questionable, the option of no treatment should also be discussed.
- Information overload: There are circumstances in which too much information may prevent the parent or guardian from making a decision. The parent or guardian may become overwhelmed with the options or potential adverse consequences and be unable to make any decision. Thus, the health care provider is not obliged to detail every statistically possible complication, even though highly unlikely (e.g., the possibility of exsanguination from an umbilical line in an infant with normal coagulation status).
- The limits of medical confidentiality: Current legal, medical, policy, and social considerations suggest that an informed consent contract with a parent or guardian needs to be defined in terms of relative rather than absolute confidentiality. Local, state, or federal regulations and laws may permit or require the reporting of medical and other information to third parties (e.g., state and federal law enforcement agencies, state and federal health agencies, insurance companies).
- The duty to warn and protect: A health care professional may have an ethical or legal duty to warn identifiable individual(s) that they have been or may have been exposed to disease or violence (contact tracing). There are some states that forbid physicians from notifying the sexual partners of patients infected with the human immunodeficiency virus (HIV). However, although the health care provider cannot force the infant's mother to inform the infant's biological father that she is infected with HIV, the health care provider can inform the infant's father that the infant is infected, regardless of whether the parents are married, especially if the father will be involved in raising the child and therefore needs to know about universal precautions.
- Duty to impute consent: There is a long-standing general medical principle that informed consent may be imputed to an unconscious accident victim who has a life-threatening condition that requires surgery. In an emergency, such rational behavior can also be imputed to the "absent parent†of an infant. When time allows, prior consultation with the hospital administration and/or hospital attorney is recommended in matters involving infants whose parents or guardians are not available or who are unable to make the necessary acute or long-term medical decisions concerning the infant. In such instances, the hospital may be forced to petition the court to appoint a legal guardian. The court may accept its historic role of parens patriae (substitute parent) and make such an appointment (22).
- Court petitions for guardianship: The court can be petitioned for the appointment of a temporary legal guardian, if the parents are unavailable or unwilling to consent to a routine medical treatment such as a life-saving blood transfusion (even if the refusal is based on sincere religious convictions such as those held by Jehovah's Witnesses [22]). If the parents are unavailable and a reasonable attempt has been made to contact them, then continuing to withhold the emergency treatment because of the failure to obtain their consent may be a basis for malpractice liability. Involvement of hospital administrative personnel and legal consultation may be advisable in such circumstances.

- All parties must understand the basic elements of informed consent as it pertains to neonates (13,14,16,17 and 18,20,21,22 and 23
 - A clear and easily understandable description of the diagnosis, the procedure, and an explanation of why the procedure is necessary for the treatment of the neonate and what may occur if the procedure is not performed.
 - A clear and easily understandable description of the reasonably foreseeable risks or discomforts.
 - A clear and easily understandable description of the benefits to the neonate. In the case of participation in a research protocol, the informed consent must contain a statement that there may be no benefit to the infant or the family.
 - A disclosure of appropriate alternative procedures or course(s) of treatment, if any, that might be advantageous to the neonate. This should include a description of what will happen to the neonate if the procedure is not performed.
 - A statement describing the extent, if any, to which the records will be protected and whether the research subjects or their parent or guardian can be identified from the medical records during or after the conclusion of the research. The statement should also indicate that research records may be inspected by the FDA and other federal agencies.
 - A statement defining who can be contacted for additional information about the treatment procedure(s).
 - The signature of the health care provider who has discussed the contents of the informed consent with the parent or guardian and the signature of the parent or guardian, dated and usually witnessed by at least one person.
 - In addition to a signed informed consent form, the author recommends that the health care provider write a note in the patient's chart indicating that the critical elements contained in the informed consent were explained to the parent or guardian. This note should also include the name(s) of those who signed and witnessed the consent. The most important aspect of an informed consent is the process of communication; the hospital chart note is simply a record that such communication has occurred. It documents the major issues that have been discussed. The more significant the illness, the greater the risks of the procedure or not performing the procedure; the more information that needs to be conveyed to the treatment decision maker, the greater is the amount of detail that should appear in the medical chart.
 - In instances in which written consent cannot be directly obtained, consent may be obtained by telephone. In such an instance, a copy of the consent form, co-signed by any witness(es) (at least one) who heard the explanation and the consent being given, should be legally sufficient. A separate note in the patient's chart should also document the parties involved in the informed consent process and the general content of the material provided to the parent or guardian during the telephone conversation.
- The individual who provides the necessary details of the planned procedure must be competent to communicate the necessary information adequately to all concerned.
 - The health care provider who is going to perform or direct the therapeutic intervention is in the best position to obtain informed consent.
 - Consent can be obtained by anyone with the necessary training to understand and explain the details of the procedure and the relevant risks.
 - Allocation of scarce and costly medical resources has become a function of availability of the resources as well as financial accessibility to those resources. Third party payers such as insurance companies and health maintenance organizations (HMOs) have altered the manner in which medical treatment decisions are perceived to be made and actually are made (24,25). Under the "managed care†system, an individual or family may wish a particular procedure to be performed (e.g., the placement of a gastrostomy tube to facilitate home or institutional care), but the procedure will not be performed unless it is either approved by the third-party payer or, if disapproved, the family members or others agree to pay for it. Thus, the treating physician may also be required to obtain "informed consent†from the

third-party payer as health care advocate for the patient. Thus, the individual responsible for obtaining consent for a procedure must be capable of gaining the trust of those to whom he presents information.

D. Policy and Regulatory Issues

• Medical records

The U.S. federal government has enacted the Health Insurance Portability and Accountability Act (HIPPA), which is administered, in large part, by the Department of Health and Human Services (DHHS), and specifically recognizes that the manner in which medical records are kept has evolved from handwritten notes maintained in filing cabinets in individual practitioners' offices to electronic databases that can be instantaneously shared with health care providers, vendors of health services, insurance companies, and employers. Patient privacy issues have become a national concern and appear to require national legislation and enforcement to ensure adequate protection. Disclosure of identifiable information, it is recommended, should be limited to the \hat{a} eminimum necessary to accomplish the purpose of the disclosure and should be used within an organization only for the purposes for which the information was collected (26). \hat{a} However, there is a need for a pragmatic balance; health information has to be accessible to public health and emergency medical services.

Medical records pertaining to the care and treatment of infants present several unique problems. The patient is not competent to provide informed consent or restrict access to the records. Guardians or parents may not be in agreement as to the nature and extent of the treatment or who should have access to medical records. Regardless of who has the legal responsibility for providing that authorization for access, treating health care providers, in general, are presumed to have access to their patient's records. A requirement for a signed consent for a routine referral can impair care by delaying consultation and referral. Thus, a physician may refer a patient to another physician (at another medical facility, including a hospital or HMO) without a specific requirement for a release of medical information form.

The Uniform Health Care Information Act permits disclosure $\hat{a} \in \infty$ to a person who is providing health care to the patient (27). $\hat{a} \in Of$ note, diagnostic interpretation of radiographic and other studies, through the use of telemedicine consultations, would appear to be covered under any statutes designed to protect medical confidentiality without compromising medical care.

Telemedicine and in-hospital consultations are conceptually similar.

The Health Insurance Portability and Accountability Act (HIPAA) of 1996 (28), required the Department of Health and Human Services (HHS) to put forth national standards for electronic health care transactions in such a manner as to prevent the erosion of the privacy of health information. However, the Law and the associated Regulations are not designed to impede the transfer of critical medical information via any media (verbal, written, or electronic) in the case of a serious or life-threatening condition. In other words, the law does not contemplate or require a delay in life-saving or other significant treatment because all the paperwork has not been properly completed. Some States have enacted "patient bill of rights†legislation to protect the rights and needs of those who may not be able to protect themselves. If the state law is more restrictive than HIPAA, it will have precedence. When it comes to interpreting state and federal laws and regulations and guidelines, it is advisable to contact the hospital legal department.

• Medical confidentiality

Patient-identifiable information should not be disclosed except as authorized by the patient or his or her guardian or as explicitly required by law. Security measures should be implemented to protect patient information against reasonably anticipated threats. Locking doors and cabinets at the end of a business day, limiting access to those who actually need to know about a given patient, and not discussing a patient by name or other identifiable characteristic in a public place continue to be part of the common-sense approach to medical confidentiality.

While disclosures of identifiable information should be limited to the minimum necessary to accomplish the purpose of the disclosure, restricting information excessively may interfere with the diagnosis, treatment, and management of the patient. Parents and guardians should understand that, in a hospital or medical center setting, ready access to patient information is, of necessity, less restricted than it is in an individual clinician's office. For example, professional staff, unknown to the family, may have access to medical records for purposes of consultation or quality assurance. In addition, the tracking of medical pharmaceuticals and medical devices, legislated and required by the Federal Food, Drug and Cosmetic Act and enforced by the FDA (29), requires that physicians report information \hat{a} . Sometimes including patient identities \hat{a} ?"to device manufacturers (30). Does the failure to provide adequate information risk compromising patient care? This is the primary question to consider in making a decision to withhold or communicate critical and sensitive medical information.

E. Unresolved Problems

- When emotional stability, maturity, or sobriety (intoxication, regardless of the nature of the psychoactive substance) is an issue in obtaining informed consent for treatment, the physician has a duty to inform a third party (the hospital legal department and the hospital department of social services) and to document his or her actions in the medical record.
- Generally, if the parents are not legally married, the informed consent of the mother is considered more substantive than that of the father. Grandparents may advise the neonate's mother, but they cannot overrule her decision unless they have legal custody of the infant. When the parents are married and they are in conflict as to the course of action to be taken, both parents' consent should be obtained before proceeding. If there is no agreement, and the issue is substantive, the hospital attorney should be consulted.
- A nonpregnant teenager, living at home and attending school, is not considered emancipated. She may not have the right, under law, to make the ultimate decisions about some aspects of her own medical care. A teenager becomes emancipated when she reaches the age of majority (in most states, age 18 years), is a member of the armed forces, is living away from home and supporting herself economically, or can otherwise demonstrate financial independence from her family to the court. However, regardless of age and financial independence, once her child is born, she has the legal right, in almost all states, to make medical decisions for her child. Thus, a 13- or 14-year-old unwed mother usually has the ultimate legal and moral responsibility for the care of her child unless the court is successfully petitioned for the transfer of custody (including medical decision-making authority) to a third party such as a relative or other court-appointed guardian.

References

1. Darling v. Charleston Community Memorial Hospital, 33 Ill.2d 326, 211 N.E.2d (1965); cert. denied, 383 U.S. 946 (1966).

- 2. 51 Fed. Reg. 22,010 (June 17, 1986).
- 3. 42 C.F.R. 403 et seq. (2004) (Medicare Conditions of Participation for Hospitals).

4. Canterbury v. Spence, 464 F.2d 772 (D.C. Cir. 1972), cert. denied, 409 U.S. 1064 (1972).

5. Cooper v. Roberts, 286 A.2d 647, 650 (1971).

6. Davis v. Wyeth, 399 F.2d 121 (9th Cir. 1968).

7. Getchell v. Mansfield, 260 Or. 174, 489 P.2d 953 (1971).

8. Levine RJ. Informed consent: some challenges to the universal validity of the western model. Law Med Health Care. 1991;19:207.

9. Levine RJ. Ethics and Regulation of Clinical Research. 2nd ed. Baltimore: Urban & Schwarzenberg; 1986.

10. Schloendorf v. Society of New York Hospital, 149 A.D. 415, 133 N.Y.S. 1143 (1912), affirmed Schloendorf v. Society of New York Hospital, 211 N.Y. 125, 105 N.E. 92 (1914).

11. Ingelfinger F. Informed (but uneducated) consent. N Engl J Med. 1972;287:465.

12. Zebarth v. Swedish Hospital Medical Center, 81 Wash.2d 12, 499 P.2d 1 (1972).

13. Keeton WP, Dobbs DB, Keeton RE, et al., eds. Prosser & Keeton on Torts. 5th ed. St. Paul, MN: West; 1984:114.

14. Ginzburg HM, MacDonald MG. Law quality assurance, and risk management in the practice of neonatology. In: MacDonald MG, Mullett MD, Sechia MMK, eds. Neonatology: Pathophysiology and Management. 6th ed. Philadelphia: Lippincott Williams & Wilkins; 2005:89–110.

15. Markowitzâ€". Arizona Parks Board, 146 Ariz. 352, 706 P.2d 365 (1985).

16. Ginzburg HM. Protection of research subjects in clinical research. In: Vevaina JR, Bone RC, Kassoff E, eds. Legal Aspects of Medicine. New York: Springer-Verlag; 1989:51–60.

17. 45 C.F.R. part 46.

18. 20 C.F.R. part 50 (Protection of Research Subjects) and part 56 (Institutional Review Boards).

19. Division of Human Subject Protections, Office of Protections from Research Risks. Findings and required actions regarding investigation of noncompliance with HHS regulations for the protection of human subjects involving the National Institutes of Health Intramural Research Program, July 3, 1991.

20. 45 C.F.R. 46.201 et seq.

21. 45 C.F.R. 46.401 et seq.

22. Application of President and Directors of Georgetown College, 331 F.2d 1000, rehearing denied, Application of President and Directors of Georgetown College, 331 F.2d 1010, and cert. denied, Jones v. President and Directors of Georgetown College, Inc., 377 U.S. 978 (1964).

23. Reiser SJ, Dyck AJ, Curran WJ. Ethics in Medicine: Historical Perspectives and Contemporary Concerns. Cambridge: MIT Press; 1977.

24. Berwick DM. Part 5: payment by capitation and the quality of care. N Engl J Med. 1996;335:1227.

25. Pegram v. Herdrich, 154 F.4d 362, reversed (98-1949).

26. Recommendations of the Secretary of Health and Human Services, Pusuant to Section 264 of the Health Insurance Portability and Accountability Act of 1996.

27. 45 CFR Parts 160–164.

28. Public Law 104-191, specifically, 45 CFR Part 160 and Subparts A and E of Part 164.

29. 21 U.S.C. §Â§301 et seq., Federal Food, Drug, and Cosmetic Act.

30. 21 U.S.C. §360i, Federal Food, Drug, and Cosmetic Act.

2-Maintenance of Thermal Homeostasis

Dora C. Rioja-Mazza

A. Definitions

- Homeostasis: Fundamental mechanism whereby living things regulate their internal environment within tolerable limits, thus keeping a dynamic equilibrium and maintaining a stable, constant condition. From the Greek homeo (same, like) and stasis (stable state) (1).
- Normal temperature: The core body temperature is maintained by the term infant within the range of 36.5ŰC to 37.5ŰC, and the skin temperature, from 0.5ŰC to 1.0ŰC lower (2).
- Thermoneutral zone: The range of ambient temperature required for the infant (for each gestational age and weight) to keep a normal body temperature (core body temperature from 36.5ŰC to 37.5ŰC) and a minimal basal metabolic rate (2,3 and 4).
- Thermoregulation: Mechanisms by which the infant tries to balance heat production and heat loss to accommodate the thermal environment (5,6 and 7).
- Cold stress: The infant senses heat loss as a stress and responds with increased heat production and peripheral vasoconstriction, with centralization of circulation, in an effort to maintain the core temperature (8).
- Hypothermia: Heat losses exceed heat production, dropping the infant's temperature below normal range (9).
 - Mild hypothermia (cold stress): $36.0\hat{A}^{\circ}C$ to $36.4\hat{A}^{\circ}C$ ($96.8\hat{A}^{\circ}F$ to $97.5\hat{A}^{\circ}F$)
 - Moderate hypothermia: $32.0\hat{A}^{\circ}C$ to $35.9\hat{A}^{\circ}C$ ($89.6\hat{A}^{\circ}F$ to $96.6\hat{A}^{\circ}F$)
 - Severe hypothermia: below $32\hat{A}^{\circ}C$ (89.6 $\hat{A}^{\circ}F$)
- Hyperthermia: An increase in the infant's temperature to above 37.5ŰC (99.5ŰF), due to a warm environment. Hyperthermia is less common than hypothermia but is equally dangerous. Clinically, it may be difficult to distinguish hyperthermia from fever (infectious origin); therefore, always consider both causes in any increase in temperature (9).

B. Background

- Effects of hypothermia:
 - Hypothermia may have severe consequences in newborn infants and may even lead to death (10,11).
 - \circ $\;$ Peripheral vaso constriction: acrocyanosis, paleness, and coldness to touch
 - Respiratory distress, apnea, and bradycardia (12,13)
 - Depletion of caloric reserves and hypoglycemia, causing shift to anaerobic metabolism and lactic acid production (14,15)
 - Increased oxygen consumption and metabolic demands result in metabolic acidosisâ€"a strong pulmonary vasoconstrictor inducing hypoxemia and central cyanosis (16,17).
 - Mobilization of norepinephrine and free fatty acids. Norepinephrine release promotes pulmonary hypertension and pulmonary ventilation–perfusion mismatch (18).
 - Risk of kernicterus at low levels of serum bilirubin (19)
 - Poor weight gain with chronic hypothermia (20)
 - Controlled hypothermia may have a neuroprotective effect in term and near-term infants with moderate to severe hypoxic ischemic encephalopathy (21,22).
- Effects of hyperthermia or overheating (9)
 - Peripheral vasodilatation: The skin is hot, the extremities are red, and the face is flushed. Diaphoresis present in full-term infants. Skin temperature is higher than core temperature.
 - Apnea, tachypnea
 - Tachycardia and hypotension

- The infant assumes a spread-eagle posture.
- Hyperactivity and irritability: The infant becomes restless and cries, then feeds poorly, with lethargy and hypotonia.
- If hyperthermia is severe, shock, seizures, and coma may occur.
- If the increase in temperature is due to hypermetabolism (infection), paleness, vasoconstriction, cool extremities, and core temperature higher than skin temperature may be noted.
- Factors affecting heat loss
 - o Infant
 - Large surface area relative to body mass
 - Relatively large head with highly vascular fontanelle
 - Skin maturation/thickness, epidermal barrier functionally mature at 32 to 34 weeks. Transepidermal water loss may be 10 to 15 times greater in preterm infants of 25 weeks' gestation (4).
 - Decreased stores of subcutaneous fat and brown adipose tissue in more premature infants (7)
 - Inability to signal discomfort or trigger heat production (shivering) (7)
 - Environment (3,4)
 - Physical contact with cold or warm objects (conduction)
 - Radiant heat loss or gain from proximity to hot or cold objects (radiation)
 - Wet or exposed body surfaces (evaporation)
 - Air currents in nursery or in incubator fan (convection)
 - Excessive or insufficient coverings or clothing
 - \circ Other factors
 - Metabolic demands of disease: asphyxia, respiratory distress, sepsis (11)
 - Pharmacologic agents, e.g., vasodilating drugs, maternal analgesics, and unwarmed IV infusions, including blood products
 - Medical stability of infant prior to procedure
 - Thermogenic response matures with increase in postconception age (4)

C. Indications

- Maintenance of thermal homeostasis is necessary at all times, but particular attention should be paid when the neonate is undergoing diagnostic or therapeutic procedures.
- Avoids increase in insensible water loss (IWL) and improves caloric utilization.

D. Equipment, Techniques, and Complications

- Prevention of heat loss in the delivery room
 - Provide a warm environment, room temperature >28.0 $\hat{A}^{\circ}C$; place infant on a radiant warmer, dry the skin with prewarmed blankets (9,14).
 - Use occlusive plastic blankets/bags (10,23).
 - Polyethylene bags (20 cm x 50 cm) prevent evaporative heat loss in infants <28 weeks' gestation. Their diathermancy allows transmission of radiant heat to the infant. Immediately after delivery, open the bag under the radiant warmer; wrap the wet infant's body from the shoulders down, and dry only the head. After stabilization, remove the wrap after the infant is inside a humidified incubator (23).
 - Environment: Maintains temperature and reduces IWL by 25% (23,24)
 - Access: Allows neonatal resuscitation (secure airway, intubation, and chest compressions), but vascular access is limited
 - Asepsis: Limited by access
 - Precautions: Record core temperature every 5 to 10 minutes until infant is stable.

- Complications: Hyperthermia, skin maceration, risk of infection
- Stockinette caps are not effective in reducing heat loss in term infants in the delivery room; there is insufficient evidence in preterm infants (10,24). Wool head coverings may reduce or prevent heat loss in term infants in the delivery room (24,25).
- Prevention of heat loss in the neonatal intensive care unit (NICU)
 - Rigid plastic heat shields (heat shielding)
 - Environment: Reduces IWL by 25% (26)
 - Access: Very limited
 - Asepsis: Limited by access
 - Precautions: Avoid direct skin contact.
 - Complications: Hyperthermia, skin maceration, risk of infection
 - Semiocclusive artificial skin (epidermal barrier protection) (27)
 - Tegaderm (3M, St. Paul, MN, USA). Semipermeable polyurethane membrane; application to extremely low-birthweight (ELBW <1,000 g) infants shortly after birth decreases fluid imbalance, allows electronic monitoring through membranes, and avoids adhesive injuries
 - Environment: Reduces IWL and improves skin care
 - Access: Limited in the area covered
 - Asepsis: Needs to be removed completely before any asepsis
 - Precautions: Thermal control inefficient for <1-kg infants
 - Complications: Risk of infection
 - Petroleum-based preservative-free ointments (epidermal barrier protection)
 - Aquaphor Original Emollient Ointment (Beiersdorf, Norwalk, CT, USA)
 - Use of this type of ointment has been discouraged in extremely low-birthweight infants (<1,000 g) because of evidence showing increase in nosocomial bacterial sepsis, caused mainly by coagulase-negative Staphylococci (28,29).



FIG. 2.1. All aspects of homeostasis are maintained during a procedure by use of an extra heat source, swaddling, stockinette cap, comfortable position, and sucrose/analgesia pacifier.

- Mechanical devices to maintain temperature
 - Thermal resistor (thermistor): A probe placed on the anterior abdominal wall or interscapular area. Used to servocontrol incubator/radiant warmer to keep infant temperature between 36.0ŰC and 36.5ŰC (4,30)
 - Convection-warmed incubator (Fig. 2.1)
 - Environment: Creates a microclimate for each infant. Infant servocontrol (ISC) triggered by skin or air temperature; temperature can also be set manually. Double plastic walls, insulated mattress, and forced-heated/humidified air minimize IWL and maintain temperature.
 - Access: Impeded by portholes, especially when working with assistants. Improved with new incubators/warmers to allow better access [e.g., Giraffe Omnibed neonatal care station (GE Medical Systems, Waukesha, WI, USA)]
 - Asepsis: Impossible to maintain wide sterile field and infant position
 - Precautions: Take infant's temperature before and after procedure. Use ISC and ensure that thermistor remains in place. Add extra heat source (heat lamp) for unstable infants or stressful procedures. Clinical deterioration may require lifting the protective shield.
 - Complications: Hyperthermia, hypothermia, unexpected break of aseptic field
 - Radiant warmed bed: For unstable infants (30)
 - Environment: Increases IWL by 50% in small preterm infants.
 - Access: Unimpeded access to infants receiving intensive care
 - Asepsis: Ability to maintain infant position and wide sterile field; also allows assistants to participate
 - Precautions: Keep infant 80 to 90 cm from radiant heat. For premature infants, heat shielding must be added. Increase fluid infusions. Record temperature every 5 to 10 minutes or use continuous monitor. To avoid burns, do not place oily substances on infant's skin.
 - Complications: Hyperthermia and dehydration
 - Heat lamp: As an extra heat source (30)
 - Environment: Increased IWL
 - Access: Limited by other equipment used (open incubator, bassinette walls)
 - Asepsis: May be affected by limited access
 - Precautions: Record temperature every 5 to 10 minutes or use continuous monitor. To avoid burns, do not place oily substances on infant's skin. Avoid heating incubator thermometer; apply manual temperature control (33°C to 35°C) when using open incubator. Keep infant approximately 60 to 90 cm from lamp bulb, and cover infant's eyes and genitals to protect from the light.
 - Complications: Cooling or overheating of isolette due to failure to detach the thermistor from infant; dehydration
 - Warming mattress: Extra heat source, for transport or radiology procedures (e.g., MRI). Effective in preventing and treating hypothermia in premature infants (<1,500 g) in the delivery room (10,24)
 - Environment: Heating through conduction, reduces heat requirements and IWL
 - Heated water-filled mattress (keep at 37.0ŰC)
 - Exothermic crystallization of sodium acetate mattress (Transwarmer Infant Transport Mattress, Prism Technologies, San Antonio, TX, USA) with a postactivation temperature of 39.0ŰC ± 1.0ŰC
 - Access: Limited only by other equipment used
 - Asepsis: Limited only by other equipment used
 - Precautions: Record temperature every 10 to 20 minutes or use ISC continuous monitor
 - Complications: Hypothermia, hyperthermia, burns

E. Special Circumstances/Considerations

- Regulate room temperature to one optimal for infant $(28\hat{A}^{\circ}C \text{ to } 30\hat{A}^{\circ}C)$ (9).
- Prewarm all heating units, including radiant warmers and incubators
- Remember that very low-birthweight preterm infants, and infants during the immediate newborn adaptation period, are more vulnerable to hypothermia and IWL.
- For transport outside of the NICU, use heated, battery-operated transport double-walled incubator.
- Plug incubator into wall outlet during procedure to allow battery to charge.
- Be aware that anesthesia may inhibit the infant's thermoregulatory capabilities.
- Warm all anesthetic and respiratory gases to body temperature and humidify.
- Gastrochisis/omphalocele: These abdominal wall defects increase risk of heat loss, fluid imbalance, and visceral damage. The infant may be placed in a "bowel bag†from the torso down, or the entire abdomen may be wrapped in clean, clear plastic wrap. Avoid visceral ischemia by keeping intestines directly above the abdominal wall defect or keep the infant in right lateral decubitus position (31).
- Neural tube defects. Keep the infant in prone position, cover the lesion with sterile gauze (soaked in warmed sterile saline), then wrap the trunk circumferentially with a dry gauze, finally covering the dry gauze with plastic wrap to minimize insensible water losses and prevent hypothermia (32).

References

1. Stedman's Medical Dictionary. 27th ed. Baltimore: Lippincott Williams & Wilkins; 2000.

2. Hey E. Thermal neutrality. Br Med Bull. 1975;31:69–74.

3. LeBlanc M. Relative efficacy of an incubator and an open warmer in producing thermoneutrality for the small premature infant. Pediatrics. 1982;69:439–445.

4. Dollberg S, Hoath SB. Temperature regulation in preterm infants: role of the skin-environment interface. Neoreviews. 2001;2:282–290.

5. Silverman WA, Zamelis A, Sinclair JC. Temperature regulation in the newborn infant. N Engl J Med. 1966;274:146–148.

6. Brück K. Temperature regulation in newborn infant. Biol Neonate. 1996;3:65–119.

7. Asakura H. Fetal and neonatal thermoregulation. J Nippon Med Sch. 2004;71:360–370.

8. Lyon AJ. Temperature control in very low birthweight infants during first five days of life. Arch Dis Child Fetal Neonatal Ed. 1997;76(1):F47–F50.

 Department of Reproductive Health and Research (RHR), World Health Organization. Thermal Protection of the Newborn: A Practical Guide. Geneva, Switzerland: World Health Organization, 1997.
McCall EM, Alderdice FA, Halliday HL, et al. Interventions to prevent hypothermia at birth in preterm and/or low birthweight babies. Cochrane Database Syst Rev. 2005;(1): CD004210.

11. Costeloe K, Hennessy E, Gibson AT, et al. The EPICure study: outcomes to discharge from hospital for infants born at the threshold of viability. Pediatrics. 2000;106:659–671.

12. Thoresen M, Whitelaw A. Cardiovascular changes during mild therapeutic hypothermia and rewarming in infants with hypoxic-ischemic encephalopathy. Pediatrics. 2000;106: 92–99.

13. Gebauer CM, Knuepfer M, Robel-Tillig E, et al. Hemodynamics among neonates with hypoxic-ischemic encephalopathy during whole-body hypothermia and passive rewarming. Pediatrics. 2006;117:843–850.

14. The International Liaison Committee on Resuscitation (ILCOR) consensus on science with treatment recommendations for pediatric and neonatal patients: neonatal resuscitation. Pediatrics. 2006;117:e978–e988.

15. Doctor BA, O'Riordan MA, Kirchner HL, et al. Perinatal correlates and neonatal outcomes of small for gestational age infants born at term gestation. Am J Obstet Gynecol. 2001;185: 652–659.

16. Hassan IA, Wickramasinghe YA, Spencer SA. Effect of limb cooling on peripheral and global oxygen consumption in neonates. Arch Dis Child Fetal Neonatal Ed. 2003,88: F139–F142.

17. Marks KH, Lee CA, Bolan CD Jr, et al. Oxygen consumption and temperature control of premature infants in a double-wall incubator. Pediatrics. 1981;68(1):93–98.

18. Endo M, Hata M, Saiki Y, et al. Hypoxia and cold stress on pulmonary venous obstruction. Pediatr Cardiol. 2001;22: 292–296.

19. Ritter DA, Kenny JD, Norton HJ, et al. A prospective study of free bilirubin and other risk factors in the development of kernicterus in premature infants. Pediatrics. 1982;69:260–266.

20. Glass L, Silverman WA, Sinclair JC. Effects of the thermal environment on cold resistance and growth of small infants after the first week of life. Pediatrics. 1968;41:1033–1046.

21. Shankaran S, Laptook AR, Ehrenkranz RA, et al., for the National Institute of Child Health and Human Development Neonatal Research Network. Whole-body hypothermia for neonates with hypoxic-ischemic encephalopathy. N Engl J Med. 2005;353:1574–1584.

22. Higgins RD, Raju TN, Perlman J, et al. Hypothermia and perinatal asphyxia: executive summary of the National Institute of Child Health and Human Development workshop. J Pediatr. 2006;148:170–175.

23. Vohra S, Roberts RS, Zhang B, et al. Heat loss prevention (HeLP) in the delivery room: a randomized controlled trial of polyethylene occlusive skin wrapping in very preterm infants. J Pediatr.

2004;145:750–753.

24. Watkinson M. Temperature control of premature infants in the delivery room. Clin Perinatol. 2006;33:43–53.

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25. Lang N, Bromiker R, Arad I. The effect of wool vs cotton head covering and length of stay with the mother following delivery on infant temperature. Int J Nurs Stud. 2004;41:843–846.

26. Symonds ME, Lomax MA. Maternal and environmental influences on thermoregulation in the neonate. Proc Nutr Soc. 1992;51:165–172.

27. Bhandari V, Brodsky N, Porat R. Improved outcome of extremely low birth weight infants with Tagaderm application to skin. J Perinatol. 2005;25:276–281.

28. Conner JM, Soll RF, Edwards WH. Topical ointment for preventing infection in preterm infants. Cochrane Database Syst Rev. 2004;(1):CD001150.

29. Edwards WH, Conner JM, Soll RF; Vermont Oxford Network Neonatal Skin Care Study Group. The effect of prophylactic ointment therapy on nosocomial sepsis rates and skin integrity in infants with birth weights of 501 to 1000 g. Pediatrics. 2004;113:1195–1203.

30. Korones S. An encapsulated history of thermoregulation in the neonate. Neoreviews. 2004;5:78–85. 31. Sheldon RE. The bowel bag: a sterile, transportable method for warming infants with skin defects. Pediatrics. 1974;53(2): 267–269.

32. Das UG, Leuthner SR. Preparing the neonate for transport. Pediatr Clin North Am. 2004;51:581–598.

3-Methods of Restraint

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Physical restraints are required for proper positioning for certain procedures. Infants may also need to be restrained to prevent accidental injury or interference with treatment (i.e., removal of feeding tubes, catheters, etc.). Select the least restrictive but most appropriate restraint for the individual patient. A. Definitions

• Physical Restraint: "any device, garment, material, or object that restricts a person's freedom of movement or access to one's body†(1)

B. Indications Restraints should be utilized:

- When procedures require proper positioning to maintain asepsis and facilitate access to patient (IV placement, lumbar punctures, radiographs, etc.) (1)
- To reduce the risk of interference with treatment (removal of feeding tubes, IV access, mechanical ventilation, etc.) (2)
- To prevent accidental injury

C. Contraindications

Restraints should not be utilized:

- When close observation of the patient could protect against potential injury or potential interference with treatment (3)
- When a change in treatment or medication regimen could protect against potential injury or interference with treatment (3)
- When modification of the patient's environment (decreased stimuli, appropriate developmental positioning, reduced noise) could protect against potential injury or interference with treatment (3)
- When use of a restraint could compromise patient care, procedures, or emergency access (1)

D. Techniques

Restraints for Procedures/Positioning

- Mummy Restraint
 - Purpose: Safe temporary method for restraining young children for treatment or examination; allows for unimpeded access to head and scalp; individual extremities can be released for access for examination or treatment (1)
 - Equipment:
 - Commercially available restraint ("papoose boardâ€) for larger infants

OR

- Clean blanket or small sheet
- Safety pins or other device for securing final blanket fold
- \circ Procedure (1):
 - Open blanket or sheet.
 - Fold one corner toward the center.
 - Place infant on blanket with shoulders at fold and feet toward opposite corner (Fig. 3.1 A).
 - With infant's right arm flexed and midline, tuck right side of blanket across trunk and under left side of body (Fig. 3.1 B).
 - Fold lower corner up toward head and tuck under left shoulder (Fig. 3.1 C).
 - With infant's left arm flexed and midline, tuck left side of blanket across trunk and under right side of body. Be sure to secure arms under blanket (Fig. 3.1 D).
- Extremity Restraint (wrist or ankle) (Fig. 3.2.)
 - Purpose: Immobilization of one or more extremities; protects infant from interfering with or removing treatment regimens (IV access, feeding tube, mechanical ventilation, etc.)
 - \circ Equipment:
 - Commercially available restraint (sheepskin and/or foam padding) for larger infants

OR

- Roll of gauze or gauze pads
- Adhesive tape
- Safety pins or other securing device

• Procedure

- Open gauze and fold in half lengthwise to reinforce material.
- Wrap wrist or ankle with gauze at least three times to create secure restraint. Caution: Do not wrap gauze too tight; this might interfere with distal circulation.
- Use adhesive tape to ensure that gauze does not unravel.
- Secure restraint to mattress, blanket, or light sandbag with safety pin.



FIG. 3.1. A: Mummy restraint: steps (1)–(3). B: Mummy restraint: step (4). C: Mummy restraint: step (5). D: Mummy restraint: step (6).

- Mitten Restraint
 - Purpose: Thumbless device to restrain or cover hand; eliminate infant's ability to grasp and possibly dislodge necessary treatment regimens (IV access, feeding tube, mechanical ventilation, etc.); prevent infant from scratching self or removing dressings, interfering with maintenance of skin integrity
 - Equipment:
 - Commercial mittens

OR

- Stockinette material (cut to fit individual infant)
- Adhesive tape
- Safety pins or other securing device (optional)



FIG. 3.2. Extremity restraint (wrist).

- \circ Procedure
 - Place infant's hand inside stockinette
 - Secure stockinette by applying tape to stockinette material and fastening around infant's wrist. Caution: Do not wrap tape too tight; this might interfere with distal circulation.
 - Tie end of stockinette in order to isolate fingers inside the stockinette material.
 - Secure restraint to mattress, blanket, or light sandbag with safety pin (optional).
- Elbow Restraint (Freedom Splint) (Fig. 3.3)
 - Purpose: Reduces ability of infant to flex elbow
 - Equipment:
 - Commercially available restraints (sheepskin and/or foam padding) for larger infants

OR

- Foam-padded armboard
- Adhesive tape
- Additional padding material, i.e., cotton balls, gauze pads
- \circ Procedure:
 - Cut four pieces of tape (appropriate size; tape should not completely encircle extremity).
 - Extend upper extremity.
 - Place armboard under elbow in order to eliminate ability to flex joint.
 - Tape extremity securely to armboard. Tape should be applied above and below elbow joint.
 - Pad bony prominences with cotton as needed.



FIG. 3.3. Elbow restraint. Restraints for Vascular Access Restraints can be used to secure IV access and prevent accidental dislodgement.

- Equipment
 - Restraint device, i.e., armboard. Armboards vary in size; a larger infant may require an armboard that is 1 to 2 cm wider than the hand/foot and extend from the proximal joint to the distal joint. However, to maintain functional position and natural curvature of the hand at rest for long-term restraint, the armboard can be shorter in length to allow for curvature of fingers around the end of the board.
 - Adhesive tape. Transparent tape is recommended for visualization of IV site especially during continuous infusion.
 - Additional padding material, i.e., cotton balls, gauze pads



FIG. 3.4. Prone positioning during procedures and at rest provides for improved breathing and sleep, lower expenditure of energy, and more stable physiologic functioning (4). Care must be taken to create positioning support of the trunk and hips.

- Procedure
 - Ensure that the infant is in a correct and functional position.
 - Rationale: Prevention of contractures and supports self-calming techniques of neonates (prone, side-lying) (Figs. 3.4 and 3.5) (4)
 - Assess skin integrity where restraint is to be applied.
 - Apply restraint board using transparent tape. Do not allow tape to encircle extremity. Three pieces of tape should sufficiently restrain extremity and allow for visualization of the tips of fingers (Fig. 3.6) or toes (Fig. 3.7). The sequence of tape allows for functional positioning of thumb and ankle.
 - Pad bony prominences and maintain natural curvature of extremities (especially the hand and fingers).



FIG. 3.5. Side-lying positioning is the best alternative to prone for procedures and sleeping. This position allows for more midline positioning of the upper and lower extremities. Nesting support increases postural stability and decreases arching of the back (4).



FIG. 3.6. Restraint for vascular accessâ€"wrist and forearm. Tape is applied in order, 1 through 3, as shown. E. Precautions

- Restraints should be a last resort after other reasonable alternatives have failed including close observation, treatment and/or medication change, modification of environment, etc. (3).
- Family education regarding the need, procedure, and time frame for the use of the restraint is required. Provide opportunity for collaboration with the family. If possible, remove the restraints when the family is visiting (1).
- For restraints during procedures, proper techniques for analgesia, sedation, and distraction (pacifier, touch, sound, etc.) are necessary in addition to the restraint.
- Weigh equipment required for restraints (i.e., armboards) prior to use. If possible, maintain a list of the weights of common restraint materials in use when weighing infants for monitoring daily growth.
- Evaluate proper use, placement, and position of restraint at least every 2 hours or sooner according to patient need, hospital policy, and regulatory agency requirement. Regulatory agencies such as the Joint Commission on Accreditation of Healthcare Organizations (JCAHO), the U.S. Food and Drug Administration (FDA), and the Centers for Medicaid and Medicare Services publish standards of medical care regarding the safe use and legal requirements for restraint implementation and maintenance (1).
- Ensure that the infant is in a proper and functional position that promotes flexion and midline positioning of upper and lower extremities.
 - Rationale: Prevention of contractures and supports self-calming techniques of neonates (prone, side-lying) (Figs. 3.4 and 3.5) (4)
- Pad bony prominences and maintain natural curvature of extremities (especially the hand and fingers)
 - Rationale: Prevents contractures and neurovascular injury; preserves skin integrity; reduces friction and pressure to skin from restraint material (1)
- When utilizing tape for securing an extremity to a board, use transparent tape when possible to allow for careful and complete assessment of the underlying skin. Do not apply tape too securely or impede circulation. Tips of the digits should remain visible for assessment.

- Restraints on the upper or lower extremities need to be assessed at least hourly for:
 - Skin Integrity, including excoriation, erythema, and edema
 - Pulses
 - Temperature
 - Color
 - Capillary Refill
 - Range of Motion (ROM) (1)
- Check for vascular circulation by inserting two fingers between infant's skin and the secured restraint (1).
 - Rationale: Constriction from a tight restraint can cause neurovascular injury and impede circulation.
- Reassess and remove restraint at least every 2 hours and with patient care (or according to individual hospital policy).
 - Rationale: Regular assessments of circulation and skin integrity and passive ROM of the joint reduce the risk of neurovascular injury and skin impairment.
- Specific assessments related to oxygenation, musculoskeletal system, and cardiorespiratory conditions need to be performed in relation to the restraint device and its usage (1).
- Observe any treatment equipment for proper positioning and patency, especially in close proximity to the restraint device (kinked IV access, dislodgement of catheters, etc.) (1).



FIG. 3.7. A: Restraint for vascular accessâ€"foot and ankle. Tape is applied in order 1 through 3, as shown. B: Foot and ankle restraint for vascular access on premature infant.

- Attach restraint to fixed location on bed (if necessary), maintaining opportunity for quick release and regular vascular checks (safety pin, secure tucking, etc.). Do not attach restraint to equipment that can be moved in such a way as to create injury or pressure to restrained joint or skin surface (crib side rails, isolette doors). Quick release allows for mobility and access in an emergency.
- Document restraint use and, if required, obtain physician order (see hospital policy).
 - Rationale: Documentation of location, time, and specific assessments maintains a record of the intervention and specific assessments that were performed. Regulatory agencies, e.g., JCAHO, FDA, and Centers for Medicaid and Medicare Services, have published standards of care addressing the documentation of patient restraints (1).
- Remove restraint at the earliest time possible.

F. Complications

- Failure of restraint resulting in self-injury and/or interference with treatment
- Neurovascular impairment (1)
- Impairment of skin integrity (i.e., pressure ulcer formation, necrosis) (1)

- Contractures or positional deformity/paralysis from prolonged immobility (1)
- Limb injury (fracture or dislocation) from movement of infant without release of secured restraint or from securing restraint to movable object (e.g., crib side rails, isolette doors) (1)
- Impairment or compromise of medical state, including oxygenation, musculoskeletal system, and cardiorespiratory conditions (1)
- Increased agitation or irritability (3)
- Extravasation injury leading to impairment of skin integrity, tissue necrosis, infection, and/or nerve and tendon damage (5)

References

1. Perry AG, Potter PA. Clinical Nursing Skills and Techniques. 6th ed. St. Louis, MO: Elsevier Mosby; 2006.

2. Ofoegbu BN, Playfor SD. The use of physical restraints of paediatric intensive care units. Pediatr Anesth. 2005;15:407–411.

3. McBeth S. Get a firmer grasp on restraints. Nursing Manage. 2004;35(10):20,22.

4. Vergara ER, Bigsby R. Developmental & Therapeutic Interventions in the NICU. Baltimore: Paul H. Brookes; 2004.

5. Ramasethu J. Prevention and management of extravasation injuries in neonates. NeoReviews. 2004;5(11):c491–c497.

4-Aseptic Preparation

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A. Definitions

- Antiseptic
 - Bactericidal or bacteriostatic substance that can be safely applied to skin
 - Not reliable as a sporicidal
 - Reduces but does not eliminate bacterial counts on the skin
 - Has an immediate effect
 - May have variable residual activity by binding to the stratum corneum of the skin
- Disinfectant
 - Chemical germicidal
 - Not sporicidal
 - Too harsh to be used on skin
- Resident flora
 - Organisms, usually of low virulence, which survive and multiply on skin and can be cultured repeatedly, e.g., Staphylococcus epidermidis
 - Cannot be completely eradicated without destroying the skin
 - Regenerate rapidly on skin when surgical gloves are worn
- Transient flora
 - Organisms that are sometimes pathogenic but do not survive and multiply on skin, e.g., gramnegative organisms such as Escherichia coli
 - \circ $\,$ Can be transmitted to patients on the hands of health care workers
 - \circ $\,$ Do not usually remain on the skin for more than 24 hours
 - Can be eradicated completely by hand washing with antiseptic solutions

B. Background

Bloodstream bacterial infection is an extremely common complication of prematurity. The majority of etiologic pathogens are nosocomial, most often transmitted by health care personnel. Use of aseptic technique is critical in reducing the number of bloodstream infections as well as in decreasing the number of contaminated blood cultures, which in turn leads to a decrease in the unnecessary use of antibiotics and the potential for antibiotic resistance. Protocols and procedures for aseptic technique in neonatal intensive care units (NICUs) are constantly being re-evaluated and updated, and hand hygiene guidelines are routinely published by the U.S. Centers for Disease Control (CDC) (1,2). Hospital managers continuously develop and update strict policies and regulations (3) as well as quality improvement projects aimed to promote adherence to aseptic technique and hand hygiene (4).

- Preparation of patient's skin and the hands of personnel prior to performing a procedure
 - \circ To remove transient flora
 - \circ $\,$ To remove and temporarily suppress most resident skin flora
- Decontamination of hands after a procedure

D. Contraindications

- Iodine solutions for preparation of skin in newborns [may cause skin and thyroid problems in high concentrations (5, 6 and 7)]
- Halogenated bisphenols (e.g., hexachlorophene) for preparation of skin in premature infants (see E7)
- Chlorhexidine for preparation of external auditory meatus

E. Precautions

- Universal (8,9)
 - Definition: A method of infection control in which all human blood and certain human body fluids are treated as if known to be contaminated with human immunodeficiency virus, hepatitis B virus, and other blood-borne pathogens
 - Indications: Reasonably anticipated risk of skin, eye, mucous membrane, or parenteral contact with blood or other potentially infectious materials, including semen, vaginal secretions, cerebrospinal fluid, synovial fluid, pleural fluid, pericardial fluid, peritoneal fluid, amniotic fluid, saliva in oral procedures, and any body fluid that is visibly contaminated with blood.
 - Major components
 - Use gloves when touching blood, body fluids, mucous membranes, or nonintact skin and when handling items or surfaces soiled with blood or body fluids.
 - Use a mask and eye protection during procedures that might generate splashing or droplets in the air.
 - Use a gown or a plastic apron when splashing of blood or body fluid is likely.
 - Wash hands carefully if they become contaminated with blood or body fluids.
 - Take extraordinary care in handling needles and other sharp objects, and dispose of them in puncture-resistant containers.
 - Exclude from patient care all personnel with exudative lesions or weeping dermatitis until these conditions have resolved.
- Recognize that no antiseptic is totally effective or without risk (Table 4.1).
- Always allow antiseptics to dry before starting procedure.
 - Drying time of at least 30 seconds is required for optimal effect (10).
 - Contamination of instruments with antiseptic is undesirable and may invalidate specimens taken for culture.

- Avoid removal of iodophore preparations prior to procedure. Removal negates the residual slow-release effect.
- After the procedure, remove iodophore from all but immediate area of procedure to prevent absorption through skin (5,6 and 7).
- Never allow antiseptic to pool under infant. Skin damage may result (16).
- Use hexachlorophene for skin preparation in newborns only as recommended by the American Academy of Pediatrics (17).
 - Use only in term infants during outbreak of Staphylococcus aureus infection.
 - Do not apply more than twice to each infant, unless application is restricted to diaper area (18).
 - Wash off solution completely.
- Reapply alcohol prior to each attempt at procedure or with any delay, as efficacy is short-lived and flora will regenerate quickly.
- Keep all antiseptics away from eyes.
- Store antiseptics in closed containers. Reusable dispensers should be thoroughly cleaned, dried, and refilled frequently. Disposable containers are available.
- Remember that gloving cannot be used as an alternative to hand washing.
 - The warm, wet skin surface under gloves offers an ideal environment for bacterial multiplication.
 - Gloves are not completely impermeable to microorganisms.
 - Latex and vinyl gloves offer comparable permeability, but vinyl gloves leak more readily.

F. Special Circumstances

- In clinical situations where traditional hand-washing facilities are unavailable, such as during patient transport, alcohol-based hand rinses, foams, or wipes may be used for hand cleaning. When alcohol solution is used, make three to five applications of 3 to 5 mL each; rub hands well until completely dry. Gloves should be used as otherwise indicated. This technique is not adequate when hands are soiled with organic matter.
- In cases of medical emergency, aseptic technique should be used as allowed by the situation, with at least use of skin preparation of the patient with an antiseptic, use of gloves, and sterile field as large as possible under the circumstances.
- Personnel suffering from allergies to antimicrobial soaps may wash thoroughly for 3 to 5 minutes with plain soap or 70% isopropanol with glycerin prior to gloving (19).
- Personnel suffering from skin cracking due to frequent use of antiseptic soaps may use skin lotions after hand washing. Products with a bacteriostatic ingredient, such as gels containing 60% ethanol, and emollients, are safe and effective in reducing skin problems (20). Containers with a flip top, rather than a screw cap, are recommended. Doebbeling et al. (21) have shown that a hand disinfection system using an antimicrobial agent (chlorhexidine) reduces the rate of nosocomial infection more effectively than one using alcohol and soap.
- Nonâ€"latex-containing gloves should be available for staff with latex allergy and to avoid allergic reactions in the patient, particularly in susceptible patients such as those with myelomeningocele (22,23).
| Considerations | Iodophor | Chlorhexidine (0.5%–4%) | Alcohol
(70%â <i>€</i> ''90%) | Iodine (1%) | Hexa-chlorophene
3%) | Chloroxylenol
(PCMX)
(0.5%â €"3.5%) |
|--|----------------------------------|---|---|--|---|---|
| 1. Indications | Hand washing
Skin preparation | Hand washing: 4%
Skin preparation: 0.5% in 70%
alcohol optimal. May be
superior to alcohol alone or
iodophor in preventing central
line infection (adult data). ^a | Skin preparation
for minor
procedures
Preparation of
external auditory
canal | Surgical hand
washing as tincture
with alcohol | Hand washing
Use limited to term
infants during
epidemics of MRSA | Hand washing
Skin and wound
disinfection |
| 2. Effective concent | ration | | | | | |
| a. Nontoxic | Hypothyroidism | Yes, ^b but local ototoxicity | Yes ^c | Hypothyroidism | CNS vacuolation | Yes ^b |
| b. Nonsensitizing | Yes | Yes | Yes | No | Yes | Low |
| c. Nonirritating | Yes | Yes | Burns in premature infants | eNo | Yes | Yes |
| 3. Mode of action | Oxidation | Cell wall disruption | Protein denaturation | Oxidation | Cell wall disruption | Cell wall disruption
and enzyme
inactivation |
| 4. Bactericidal | Yes | Yes | Yes | Yes | No | Yes |
| 5. May be used with detergent | n Yes | Yes | No | No | Yes | Neutralized by nonionic surfactants |
| 6. Persistent local action | Yes | Yes | No | Yes | Yes, but only with
repeated use; reduced
by concomitant use of
alcohol | Good |
| 7. Effective against a. Gram-positive bacteria | Yes | Yes | Yes | Yes | Yes
(bacteriostatic) | Good (better against
Streptococcus than
Staphylococcus) |

TABLE 4.1 A Comparison of Commonly Used Antiseptics (11,14,15)

b. Gram-negative	Yes	Yes	Yes	Yes	No	Fair, improved by
bacteria						EDTA
c. Spores	No	No	No	No	No	No
d. Tubercle	No	No	Yes	Yes	No	Fair
bacillus						
e. Viruses	Yes	Yes	Lipophilic viruses	Yes	Yes	Fair
			only			
f. Fungi	Yes	Yes	Yes	Yes	Yes	Fair
8. Use associated	No	Contamination with	No	No	Yes	Not effective against
with resistance		Pseudomonas and Proteus				Pseudomonas species
		species				
9. Rapid action	No	Yes (better when combined	Yes (drying time)	Yes	No	Intermediate
	Requires 4–5 min	with alcohol)				
	of scrubbing					
10. Easily	No	No	May be inactivated	l Yes	Yes	Minimal
inactivated by	Low surface tension		by nonbacterial			
extraneous organic	for good crevice and		protein			
matter	fat penetration					
MRSA, methicillin	resistant staphylococc	us aureus; CNS, central nervou	s system; EDTA, eth	ylenediaminetetra	-acetic acid.	

^aData from Maki et al. (11). ^bSkin absorption not studied in very low-birthweight infants. ^cData from Harpin and Rulter (39).

G. Technique (See Procedures DVD for Video)

A 3- to 5-minute $\hat{a} \in \hat{c}$ (vigorous washing up to the elbows) is necessary upon entering the nursery. Subsequently, a 15- to 30-second handwashing is indicated prior to and after each patient contact.

- Preparation for a minor procedure
 - Definition of a minor procedure
 - Short duration (5 to 10 minutes); noncomplex
 - Does not involve an area, such as the central nervous system (CNS), which is especially vulnerable to infection
 - Does not require skin incision
 - For example, blood drawing, placement of percutaneous peripheral venous line, bladder tap, punch-skin biopsy
 - Preparation of personnel
 - Wear cap/beard cover if hair is likely to contaminate the field.
 - Remove all jewelry from hands and arms.
 - Wash hands, wrist, forearms, and elbows using a small amount of antiseptic preparation (e.g., iodophore or chlorhexidine). Iodophore preparations appear to be equally effective when applied with disposable sponges or brushes. Vigorous scrubbing with a brush leads to skin breakdown and possible contamination and is contraindicated. Be sure to include between the fingers and the lateral surface of the fifth finger.
 - Clean nails with stick.
 - Wash/scrub hands and forearms to elbow with antiseptic for a further 2 to 3 minutes.
 - Rinse hands and forearms with running water, keeping them elevated above elbows.
 - Use towel to shut off water if knee- or foot-operated faucets are not available.
 - Dry hands with clean towel prior to drying forearms.
 - Wear gloves.
 - Preparation of patient skin
 - If necessary, cut hair in area of procedure with small scissors, taking care not to nick skin. Avoid shaving, as that may cause skin compromise and increases the risk of infection.
 - Apply antiseptic.
 - Alcohol may be used. Preparation with iodophore may be optimal, but color tends to obscure underlying vessels.
 - Apply three times in circles progressing away from procedure site.
 - Apply with some friction.
 - Allow to dry. Do not wipe off antiseptic.
 - Never touch skin after application of antiseptic and before initiation of the procedure.
 - If using alcohol, reapply it prior to every attempt at procedure.
 - Preparation for a major procedure
 - Definition of major procedure
 - Invasive or involving skin incision
 - For example, central line placement, cutdown, chest tube, lumbar puncture
 - Duration longer than 5 to 10 minutes
 - Masks, drapes, and gowns. Clothing is an important barrier to microorganisms shed into the air from the skin and mucous membranes. The pore size of gowns and masks should prevent bacterial passage even when wet [use 140 thread count or higher unless plasticized (24)]. Disposable gowns and drapes manufactured from nonwoven materials are effective in reducing infection (25). When woven reusable materials are used, they should be tightly woven and treated with a water repellent.

- Put on cap and mask.
- Clean nails and "scrub†as for minor procedure, but continue for 4 to 5 minutes.
- Rinse forearms and hands, keeping them elevated above elbows.
- Dry hands and then forearms with two sterile towels. Keep wrists and hands elevated until drying is complete.
- Put on sterile gown with the aid of an assistant (Fig. 4.1).
- Put on sterile gloves, without contaminating external surface with ungloved hand (Fig. 4.2).
 - Have assistant open packet without contaminating contents.
 - Pull gloves well over sleeve ends.
- Preparation of patient skin

С

- Prior to procedure, have assistant:
 - Wash area, if soiled, with soap and water.
 - If necessary, remove hair using small scissors, taking care not to nick skin. Do not shave the area.
- Apply antiseptic with three separate sponges. Start at center of circle, and work centrifugally to at least 5 cm outside immediate area of procedure. Alcohol (70%) should not be used. An iodophore preparation is commonly used in nurseries in the United States.
- Allow antiseptic to dry. Do not wipe off antiseptic prior to procedure.



D

FIG. 4.1. Correct technique for putting on a sterile gown. Operator is assisted into gown. A: The assistant pulls the gown up and back over the operator's shoulders by grasping the inside surface and ties the neck ties at the back of the operator's neck. B: Operator hands tip (protected with a removable cardboard tab) of sterile tie to assistant. C, D: Operator carries the tie around to the front where the operator takes tie (without the cardboard tab) and ties the gown.

H. Complications

- Dry skin with repeated use
- Hexachlorophene
 - Transcutaneous absorption with CNS vacuolation (26,27,28 and 29)
 - Possible teratogenicity when used for hand washing by pregnant staff member (30,31)
- Iodine
 - Burns
 - Allergic contact dermatitis has been reported (32).
 - Skin absorption/hypothyroidism (5,6 and 7)
 - A high incidence of transient neonatal hypothyroidism has been observed in premature infants in Europe after routine skin cleansing with iodine. The same high incidence has not been noted in North America. This difference in incidence may be due to the prior iodine status of the neonate (5).
- Iodophore
 - Burns possible when allowed to pool under infant
 - Absorption through skin reported in burn patients and neonates (6)
 - Alteration of thyroid function (5,7)
- Chlorhexidine
 - Similarity in name and preparation has led to some confusion between chlorhexidine and hexachlorophene. These compounds are different in structure and properties (Table 4.1).
 - Ototoxicity when instilled into middle ear (33)
 - Burns possible when allowed to pool under infant
 - Absorption through skin and from umbilical stump (34,35). No associated pathology was documented.
 - Contamination with gram-negative organisms has been reported, in particular, Pseudomonas and Proteus species (36,37).
 - There is no evidence that the detergent or alcohol preparations are susceptible to contamination.
- Alcohol burns in premature infants (38,39)
- Latex allergy in operators and in infants with neural tube defects (22,23).





FIG. 4.2. Correct technique for putting on sterile gloves. A: Assistant has opened outer pack, allowing removal of uncontaminated inner pack by operator. B: Correct method for lifting second glove with gloved hand to avoid contact with skin as second glove is pulled up over sleeve ends. C: Pulling first glove up over sleeve ends. The inside surface of the glove is never touched by the gloved hand. References

1. Centers for Disease Control and Prevention. Guideline for hand hygiene in health-care settings. MMWR. 2002;51:1.

2. Centers for Disease Control and Prevention. Guidelines for the prevention of intravascular catheterrelated infections. MMWR. 2002;51:1.

3. Kilbride HW, Powers R, Wirtschafter DD, et al. Evaluation and development of potentially better practices to prevent neonatal nosocomial bacteremia. Pediatrics. 2003;111:504.

4. Pittet D. Improving adherence to hand hygiene practice: a multidisciplinary approach. Emerg Inf Dis. 2001;7:234.

5. Brown RS, Bloomfield S, Bednarek FJ, et al. Routine skin cleansing with povidone–iodine is not a common cause of transient neonatal hypothyroidism in North America: a prospective controlled study. Thyroid. 1997;7:395.

6. Mitchell IM, Pollock JC, Jamieson MP, et al. Transcutaneous iodine absorption in infants undergoing cardiac operation. Ann Thorac Surg. 1991;52:1138.

7. Gordon CM, Rowitch DH, Mitchell ML, et al. Topical iodine and neonatal hypothyroidism. Arch Pediatr Adolsc Med. 1995; 149:1336.

8. Perspectives in Disease Prevention and Health Promotion Update: universal precautions for prevention of transmission of human immunodeficiency virus, hepatitis B virus, and other blood borne pathogens in health-care settings. MMWR. 1988;37:377.

9. Guidelines for prevention of transmission of human immunodeficiency and hepatitis B virus to healthcare and public safety workers. MMWR. 1989;38:3.

10. Intravenous Nurses Society. Intravenous nursing standards of practice. J Intrav Nurs. 1998;21:51. 11. Maki DG, Ringer M, Alvarado CJ. Prospective randomized trial of povidone–iodine, alcohol, and chlorhexidine for prevention of infection associated with central venous and arterial catheters. Lancet. 1991;338:339.

12. Webster J. Hand-washing in a neonatal intensive care unit: comparative effectiveness of chlorhexidine gluconate 4% w/v and triclosan 1% w/v. J Aust Coll Midwives. 1991;4:25.

13. Webster J, Faoagali JL. An in-use comparison of chlorhexidine gluconate 4% w/v, glycol-poly-siloxane plus methylcellulose and a liquid soap in a special care baby unit. J Hosp Infect. 1989;14:141.

14. Kinironos B, Mimoz O, Lafendi L, et al. Chlorhexidine versus povidone–iodine in preventing colonization of continuous epidural catheters in children: a randomized, controlled trial. Anesthesiology. 2001;94:239.

15. Herruzo-Cabrera R, Vizcaino-Alcaide MJ, Fdez-Acinero MJ. Usefulness of an alcohol solution of Nduopropenide for the surgical antisepsis of the hands compared with handwashing with iodine–povidone and chlorhexidine: clinical essay. J Surg Res. 2000;94:6.

16. Wilkinson AR, Baum JD, Keeling JW. Letter to the editor: superficial skin necrosis in babies prepared for umbilical arterial catheterization. Arch Dis Child. 1981;56:237.

17. American Academy of Pediatrics, American College of Obstetricians and Gynecologists. Guidelines for Perinatal Care. 5th ed. Elk Grove Village, IL: American Academy of Pediatrics; Washington, DC: American College of Obstetricians and Gynecologists; 2002.

18. American Academy of Pediatrics. Report of the Committee on Infectious Diseases. 19th ed. Evanston, IL: American Academy of Pediatrics; 1982:243.

19. Rotter ML. Hygienic hand disinfection. Infect Control. 1984; 5:18.

20. Newman JL, Seitz JC. Intermittent use of an antimicrobial hand gel for reducing soap-induced irritation of health care personnel. Am J Infect Control. 1990;18:194.

21. Doebbeling BN, Stanly GL, Sheetze CT, et al. Comparative efficiency of alternative hand-washing agents in reducing nosocomial infections in intensive care units. N Engl J Med. 1992;327:88.

22. Pittman T, Kiburz J, Gabriel K, et al. Latex allergy in children with spina bifida. Pediatr Neurosurg. 1995;22:96.

23. Banta JV, Bonanni C, Prebluda J. Latex anaphylaxis during spinal surgery in children with myelomeningocele. Dev Med Neurol. 1993;35:543.

24. Moylan JA, Fitzpatrick KT, Davenport KE. Reducing wound infections. Arch Surg. 1987;122:152.25. Laufman H, Eudy WW, Vandernoot AM, et al. Strike-through of moist contamination by woven and non-woven surgical materials. Ann Surg. 1975;181:857.

26. Curley A, Hawk RE, Kimbrough RD, et al. Dermal absorption of hexachlorophene in infants. Lancet. 1971;2:296.

27. Anderson JM, Cockburn F, Forfar JO, et al. Neonatal spongioform myelinopathy after restricted application of hexachlorophene disinfectant. J Clin Pathol. 1981;34:25.

28. Rossiter EJR. Hexachloropheneâ€"time to stop. Aust Paediatr J. 1980;16:236.

29. Martin-Bouyer G, Lebreton R, Toga M, et al. Outbreak of accidental hexachlorophene poisoning in France. Lancet. 1982;1:91.

30. Check W. New study shows hexachlorophene is teratogenic in humans. JAMA. 1978;240:513.

31. Halling H. Suspected link between exposure to hexachlorophene and malformed infants. Presented at the New York Academy of Science, July 1978.

32. Lee SK, Zhai H, Maibach HI. Allergic contact dermatitis from iodine preparations: a conundrum. Contact Dermatitis. 2005;52:184.

33. Brickwell PG. Sensorineural deafness following myringoplasty operations. J Laryngol Otol. 1971;85:957.

34. Agett PJ, Cooper LV, Ellis SH, et al. Percutaneous absorption of chlorhexidine in neonatal cord care. Arch Dis Child. 1981;56:878.

35. Cowan J, Ellis SH, McAinsh J. Absorption of chlorhexidine from the intact skin of newborn infants. Arch Dis Child. 1979;54:379.

36. Bassett DC, Stokes KJ, Thomas WRG. Wound infection with Pseudomonas multivorans. Lancet. 1970;1:1188.

37. Wishart MM, Riley TV. Infection with Pseudomonas maltophilia. Hospital outbreak due to contaminated disinfectant. Med J Aust. 1976;2:710.

38. Bonacci H. Letter to the editor: hazard in the nursery. N Engl J Med. 1970;282:633.

39. Harpin V, Rulter N. Percutaneous alcohol absorption and skin necrosis in a premature infant. Arch Dis Child. 1982;57:477.

5-Analgesia and Sedation in the Newborn

Nicholas J. Marsh

A. Introduction

Pain management is an integral part of compassionate neonatal medical care. Historically, barriers to adequate pain management in neonates have been related to the question of pain perception in the newborn. This question is no longer debated, and the use of analgesics for these patients is well established (1,2,3,4 and 5). However, pain management and sedation practices continue to vary among practitioners (6,7). Safe care requires careful choice and dosing of medications, appropriate monitoring, and preparedness to manage complications (4,8). This chapter offers general guidelines for the management of analgesia and sedation in newborn infants.

B. Definitions

- Analgesia: A condition in which nociceptive stimuli are perceived but are not interpreted as pain; usually accompanied by sedation without loss of consciousness (9)
- Conscious sedation: A medically controlled state of depressed consciousness that allows protective reflexes to be maintained, retains the ability to maintain a patent airway independently and continuously, and permits appropriate responses by the patient (8)
- Deep sedation: A medically controlled state of depressed consciousness or unconsciousness from which the patient is not easily aroused. It may be accompanied by a partial or complete loss of protective reflexes and includes the inability to maintain a patent airway independently and respond purposefully to stimulation (8).
- Tolerance: The ability to resist the action of a drug or the requirement for increasing doses of a drug, with time, to achieve a desired effect (10)
- Withdrawal: The development of a substance-specific syndrome that follows the cessation of, or reduction in, intake of a psychoactive substance previously used or administered regularly (11)
- Neonatal abstinence syndrome: Onset of withdrawal symptoms in neonates upon cessation of an agent associated with physical dependence (11)
- C. General Indications
 - Any condition or procedure known to be painful (4,12) (see E)
 - Physiologic indications consistent with perception of pain (1,13)
 - \circ Tachycardia
 - Tachypnea
 - Elevated blood pressure (with secondary increase in intracranial pressure)
 - Decreased arterial oxygen saturation
 - Hyperglycemia secondary to hormonal and metabolic stress responses
 - Behavioral indications consistent with perception of pain (4,5,13,14,16)
 - Simple motor responses (i.e., withdrawal of an extremity from a noxious stimulus)
 - Facial expressions (i.e., grimace)
 - Altered cry (primary method of communicating painful stimuli in infancy)
 - Agitation
 - 0

D. Specific Indications

• Analgesia

In general, the potency of analgesic treatment selected should be related directly to the anticipated or assessed level of pain (1).

- \circ Mild pain
 - Nonpharmacologic approaches (see H)
 - Local and/or topical anesthesia
 - Nonopioid analgesics (e.g., acetaminophen)
- Moderate and severe pain
 - Intravenous opioid analgesics (see E)
 - Local and/or topical anesthesia
 - Benzodiazepines (in combination with D2a; see E)
- Sedation

Sedatives may be co-administered with analgesics to enhance the anticipated benefits. Because of the escalated risks associated with deep sedation, conscious sedation should be the usual clinical endpoint.

- Benzodiazepines (see E)
- Chloral hydrate (see E)
- Nonpharmacologic approaches (see H)

E. Precautions

- Be aware, when assessing patients, that:
 - The clinical assessment of pain in the newborn is imprecise.
 - Physiologic and behavioral indicators of pain are nonspecific and may be related to many other factors.
 - Intubated neonates receiving muscle relaxants may have altered physiologic indicators and completely ablated behavioral indicators.
 - A high index of suspicion is required to identify newborn infants in pain (13).
- Be aware, when medicating patients, that:
 - There are numerous potential complications associated with analgesic and sedative agents (Table 5.1) (17,18).
 - Large inter- and intraindividual variations in response may occur (19).
 - Medications must always be titrated slowly (4).
 - Co-administration of opioids, benzodiazepines, and other sedatives may result in greatly exaggerated respiratory depressant effects, including apnea (20,21).
- Resuscitation equipment and medications should be immediately available. Be prepared to support ventilation and perform tracheal intubation if needed; respiratory depression is a common side effect of a number of analgesic agents.
- Be aware that:
 - Newborn infants who have developed tolerance to a sedative or analgesic agent, by either direct or in utero exposure, may exhibit symptoms of the neonatal abstinence syndrome upon abrupt cessation or reversal of the administered agent (20,21). For example, naloxone administered to opioid-dependent neonates may precipitate acute, severe withdrawal symptoms (11).
 - Chronic analgesic therapy with agents known to induce tolerance should be weaned gradually, with close monitoring for evidence of withdrawal symptoms (4,20,21).

- When using analgesics for a painful procedure:
 - Consider both the duration and the anticipated pain when selecting medications and methods.
 For example, short procedures with mild to moderate discomfort, such as lumbar puncture, may be best managed with topical and local anesthetics (22,23 and 24).
 - Minimize the number of painful episodes. Multiple procedures performed at the same time may avoid the need for repeated administration of analgesics (2,15).
 - Ensure that oxygen, suction, airway, resuscitation equipment, and reversal agents are readily available.
 - Follow nothing-by-mouth (NPO) guidelines for ambulatory surgery.
 - Have a nurse, or other professional not involved in the procedure, constantly monitor respirations, pulse oximetry, heart rate, and level of consciousness.
- Chloral hydrate, previously regarded as a highly safe sedative, should be used with caution in neonates (particularly premature neonates) owing to the risk of hyperbilirubinemia and accumulation of toxic metabolites (25,26 and 27). For these reasons, a single dose only is recommended.

F. Advantages and Disadvantages of Commonly Used Agents in the Newborn See Table 5.1.G. Complications See Table 5.1.H. Nonpharmacologic Approaches

- Swaddling during heel-stick procedures has been shown to reduce behavioral pain responses (28).
- Non-nutritive sucking has been demonstrated to significantly reduce crying in response to painful stimuli (29,30).
- Sucrose
 - Infants who drank 2 mL of a 12% sucrose solution prior to blood collection via heel stick cried 50% less than control infants during the same procedure (30).
 - Infants who received sucrose on a pacifier prior to and during circumcision cried significantly less than control infants (30).
 - 2 mL of 12% to 50% sucrose administered orally 2 minutes prior to the procedure is an effective neonatal analgesic with few adverse effects (31,32).

I. Contraindications

- There are no absolute contraindications to using analgesia and/or sedation when deemed clinically appropriate.
- Be aware of the potential side effects associated with the specific agent selected and take the proper precautions.

	C 4					
Agent	Category	Action	Advantages	Disadvantages	Recommended Dose	Metabolism
Acetaminophen	Oral analgesic	Inhibition of peripheral afferent pain signals	No respiratoy depression	Mild analgesia	10–20 mg/kg p.o., p.ı	. Hepatic
Fentanyl	Synthetic narcotic	Opioid receptor	Rapid onset of action	n Short duration (30–60 min)	1–5 µg/kg i.v. bolu	s Hepatic
(Sublimaze)	analgesic	agonist	(3–5 min)	Respiratory depression	0.5–1.5 µg/kg/h	
			More potent than	Hypotension	continuous infusion	
			morphine	Risk of seizures	(analgesia)	
			Reversible with	Chest wall rigidity with rapid infusion		
			narcotic antagonist	Tolerance and withdrawal	•	
Morphine	Narcotic analgesic	Opioid receptor	Longer duration than	Slower onset of action than fentanyl	20 Aµg/kg/h (ventilated	l Hepatic
		agonist	fentanyl (4 h)	(10–15 min)	patients)	
			Reversible with	Respiratory depression		
			narcotic antagonist	More adverse cardiovascular side effects		
				than fentanyl		
				Tolerance and withdrawal		
Meperidine	Narcotic analgesic	Opioid receptor	Longer duration than	Slower onset of action than fentanyl (10	0.5–1.5 mg/kg i.v. or	Hepatic
(Demerol)		agonist	fentanyl (2–4 h)	min)	1.m.	
			Reversible with	Tolerance and withdrawal		
			narcotic antagonist			
			Less respiratory			
			depression than			
		D ' (morphine			TT /
Diazepam" (Valiu	m)Benzodiazepine	Brainstem	Sedation	No analgesic effect	0.1a€ ⁴ 0.2 mg/kg 1.v.	Hepatic
	sedative/hypnotic	reticular	Music relaxation	Respiratory depression		
		formation	Rapid onset (1a€ ² 2	Hypotension With descent constants		
T a	D	Designation	IIIII) Calatian	withdrawai symptoms may occur	0.052040 1	11
Lorazepam	Benzodiazepine	Brainstem	Sedation	No analgesic effect	0.05a€ 0.1 mg/kg 1.v.	Hepatic

TABLE 5.1 Sedative and Analgesic Agents Commonly Used in the Newborn Mechanism of

(Ativan)	sedative/hypnotic	reticular	Longer duration than	Respiratory depression		
		formation	diazepam (8–12 h)	Hypotension		
		depressant		Withdrawal symptoms may occur		
Midazolam ^a	Benzodiazepine	Brainstem	Sedation	No analgesic effect	0.05–0.15 mg/kg i.v.	Hepatic
(Versed)	sedative/hypnotic	reticular	Rapid onset of action	Respiratory depression	loading dose	
		formation	(2–5 min)	Hypotension	0.16–1 µg/kg/min	
		depressant		Withdrawal symptoms may occur	continuous infusion	
Phenobarbital	Barbiturate	Nonspecific CNS	Sedation	No analgesic effect	2–10 mg/kg i.v. i.m.,	Hepatic
	sedative	depressant	Long half-life	Slower onset of action than other	p.o.	-
			Serum levels easily	barbiturates (5 min)		
			monitored	No specific antagonist		
			Reduces serum	Respiratory depression		
			bilirubin levels			
Chloral hydrate	Sedative	Central	Sedation	No analgesic effects	25–50 mg/kg p.o., p.r.	Hepatic
		depressant effect	Minimal	Gastrointestinal irritant	One administration	
			cardiorespiratory side	eMay accumulate in neonates	onlyâ€"repeated doses	
			effects	May cause myocardial depression	increase risk of	
				Associated with hyperbilirubinemia	accumulation and	
				Toxic metabolites may accumulate with	toxicity	
				repeated dosing		
EMLA cream	Topical anesthetic	Delivery of local	Painless	Not recommended in neonates <1 mo old	, Age 0–3 mo:	Hepatic,
(lidocaine 2.5%,		anesthetic to	administration, direct	t contraindicated in patients with	maximum 1 g, 10 cm^2 ,	renal
prilocaine 2.5%		dermal and	application to intact	congenital or idiopathic	≤ 1 h	
cream)		epidermal tissue	skin	methemoglobinemia or those receiving	Age 3–12 mo:	
				methemoglobininducing agents	maximum 2 g, 20 cm^2 ,	
				Systemic absorption of active ingredients	≤ 4 h	
				with formation of methemoglobinemia		
Data from refs. 1.2.	4,5,13,19,21,26					

Data from refs. 1,2,4,5,13,19,21,26 CNS, central nervous system. ^aLimited data available in newborns.

References

1. Vitali SH, Camerota AJ, Arnold JH. Anaesthesia and analgesia in the newborn. In: Macdonald MG, Mullett MD, Sechia MMK, eds. Neonatology: Pathophysiology and Management of the Newborn. 6th ed. Philadelphia: Lippincott Williams & Wilkins; 2005:1557.

2. Larsson BA. Pain management in neonates. Acta Paediatr. 1999;88:1301.

3. Anand KJS. Clinical importance of pain and stress in preterm neonates. Biol Neonate. 1998;73:1.

4. Menon G, Anand KJ, McIntosh N. Practical approach to analgesia and sedation in the neonatal intensive care unit. Semin Perinatol. 1998;22:417.

5. Stevens B. Management of painful procedures in the newborn. Curr Opin Pediatr. 1996;8:102.

6. Porter FL, Wolf CM, Gold J, et al. Pain and pain management in newborn infants: a survey of physicians and nurses. Pediatrics. 1997;100:626.

7. McLaughlin CK, Hull JG, Edwards WH, et al. Neonatal pain: a comprehensive survey of attitudes and practices. J Pain Symp Manage. 1993;8:7

8. Committee on Drugs, American Academy of Pediatrics. Guidelines for monitoring and management of pediatric patients during and after sedation for diagnostic and therapeutic procedures. Pediatrics. 1992;89:1110.

9. Stedman's Medical Dictionary. 27th ed. Baltimore: Williams & Wilkins; 2000.

10. American Psychiatric Association. Diagnostic and Statistical Manual of Mental Disorders. 4th ed. rev. Washington, DC: American Psychiatric Press; 1994.

11. Franck L, Vilardi J. Assessment and management of opioid withdrawal in ill neonates. Neonatal Networks. 1995;14:39.

12. Anand KJ, Barton BA, McIntosh N, et al. Analgesia and sedation in preterm neonates who require ventilatory support: results of the NOPAIN trial. Neonatal Outcome and Prolonged Analgesia In Neonates. Arch Pediatr Adolesc Med. 1999; 153:331.

Mathew PJ, Mathew JL. Assessment and management of pain in infants. Postgrad Med J. 2003;79:438.
 McRae ME, Rourke DA, Imperial-Perez FA, et al. Development of a research-based standard for assessment, intervention, and evaluation of pain after neonatal and pediatric cardiac surgery. Pediatr Nurs. 1997;23:263.

15. Weisman SJ, Schechter NL. The management of pain in children. Pediatr Rev. 1991;12:237.

16. Marshall RE. Neonatal pain associated with caregiving procedures. Pediatr Clin North Am. 1989;36:885.

17. Cote CJ, Karl HW, Notterman DA, et al. Adverse sedation events in pediatrics: analysis of medications used for sedation. Pediatrics. 2000;106:633.

18. Cote CJ, Notterman DA, Karl HW, et al. Adverse sedation events in pediatrics: a critical incident analysis of contributing factors. Pediatrics. 2000;105:805.

19. Jacqz-Aigrain E, Burtin P. Clinical pharmacokinetics of sedatives in neonates. Clin Pharmacokinet. 1997;36:449.

20. Caron E, Maguire DP. Current management of pain, sedation, and narcotic physical dependency of the infant on ECMO. J Perinat Neonat Nurs. 1990;4:63.

21. Notterman DA. Sedation with intravenous midazolam in the pediatric intensive care unit. Clin Pediatr. 1997;36:449.

22. Porter FL, Miller JP, Sessions CF, et al. A controlled clinical trial of local anesthesia for lumbar punctures in newborns. Pediatrics. 1991;88:663.

23. Essink-Tjebbes CM, Hekster YA, Liem KD, et al. Topical use of local anesthesia in neonates. Pharm World Sci. 1999;21: 173.

24. Taddio A, Stevens B, Craig K, et al. Efficacy and safety of lidocaine–prilocaine cream for pain during circumcision. N Engl J Med. 1997;24:1197.

25. Lambert GH, Muraskas J, Anderson CL, et al. Direct hyperbilirubinemia associated with chloral hydrate administration in the newborn. Pediatrics. 1990;86:277.

26. Reimche LD, Sankaran K, Hindmarsh KW, et al. Chloral hydrate sedation in neonates and infantsâ€"clinical and pharmacologic considerations. Dev Pharmacol Ther. 1989;12:57.

27. Mayers DJ, Hindmarsh KW, Sankaran K, et al. Chloral hydrate disposition following single-dose administration to critically ill neonates and children. Dev Pharmacol Ther. 1991;16:71.

28. Campos R. Soothing pain-elicited distress with swaddling and pacifiers in early infancy. Presented at Sixth International Conference on Infant Studies, Washington, DC, April 1988.

29. Carbajal R, Chauvet X, Coudec S, et al. Randomized trial of analgesic effects of sucrose, glucose and pacifiers in term neonates. Br Med J. 1999;319:1393.

30. Blass EM, Watt LB. Suckling and sucrose-induced analgesia in human newborns. Pain. 1999;88:611.

31. Carbajal R, Lenclen R, Gajdos V, et al. Crossover trial of analgesic efficacy of glucose and pacifier in very preterm neonates during subcutaneous injections. Pediatrics. 2002;110:389.

32. Horwitz N. Does oral sucrose reduce the pain of neonatal procedures? Arch Dis Child. 2002;87:80.

6-Temperature Monitoring

Monisha Bahri

Temperature measurement is an important part of normal newborn care. Accurate measurement is important to detect deviations from normal and also for optimal incubator and radiant warmer function.

The purpose of monitoring temperature is to maintain the infant in a thermoneutral environment zone. This is defined as the narrow range of environmental temperature in which the infant maintains a normal body temperature without increasing metabolic rate and hence oxygen consumption.

Temperature monitoring may be done intermittently or continuously. The site of measurement may be core (rectum, esophagus, or tympanic) or surface (skin, axilla). Currently, the axillary route is preferred, especially for preterm neonates. The various methods are discussed further in this chapter.

Intermittent Temperature Monitoring

A. Equipment

- Mercury-in-glass thermometer
 - o Benchmark standard
 - Determination time >3 minutes.
 - Risk of breakage/mercury poisoning from vaporized mercury (1).
 - American Academy of Pediatrics (AAP) recommendation is that mercury thermometers be removed from medical offices, medical facilities, and homes (1).
 - No longer used in neonatal units
- Electronic digital thermometers (Fig. 6.1)
 - Most widely used
 - Thermometer is a small hand-held device.
 - Temperature sensor may be a thermistor or thermocouple.
 - Temperature is sensed by the probe; the signal is then processed electronically and displayed digitally. There is an audible signal at the end of the determination time window.
 - Determination time is <45 seconds.
 - \circ Resolution is 0.1ŰC.
 - Probe-type electronic thermometers are designed to be used with disposable probe covers.
- Infrared electronic thermometry
 - Sensitive infrared sensor detects infrared energy radiation from the tympanic membrane.
 - Sensor converts the infrared signal to an electrical signal.
 - Electrical signal is then processed and displayed digitally as temperature.
 - Determination time is <2 seconds.
 - Designed to be used with disposable sensor head covers.
 - Cost-effective (2)

B. Contraindications

• Rectal measurement in very low-birthweight infants

C. Limitations

- Infrared thermometers have less correlation with glass thermometer measurements than digital disposal or electronic thermometers (2).
- Studies have failed to show adequate correlation between infrared thermometer readings and rectal or axillary measurements in newborns (3,4).
- Readings can vary from axillary site depending on environment (radiant warmer, open crib, or incubator) (5).

D. Precautions

- Probe-type electronic thermometers
 - Always use disposable probe cover.
 - Do not force probe, as perforation can occur, e.g., in rectal measurement.
- Infrared thermometers
 - Always use disposable sensor head covers.
 - Do not force sensor head into the ear canal.
 - Do not use in infants with middle ear disease.
 - Do not use in very low-birthweight infants because of inappropriate speculum size. Sensor head may not be small enough for low-birthweight infants weighing less than 1,000 g.
 - Erroneous readings may result from:
 - Not having the probe lined up with the tympanic membrane
 - The presence of heavy cerumen
 - The presence of serous otitis media (6)



FIG. 6.1. Electronic thermometers: Probe thermometer. (Courtesy of Welch Allyn, New York, USA.)

E. Technique

- Probe-type electronic thermometers
 - Apply disposable probe cover to the probe.
 - For core temperature, insert probe into the rectum (2 to 3 cm).
 - For noninvasive approximation of core temperature, place the probe in the axilla (Fig. 6.2) (7,8 and 9).
 - Hold the probe in place and wait for an audible beep before removing the probe.
 - Read temperature and return the probe to its compartment to deactivate the unit.
- Infrared thermometers
 - Apply disposable cover to the sensor head.
 - Gently insert tapered end into the ear canal.
 - While holding the unit steady, depress trigger.
 - Remove from the ear canal and read temperature.
 - Remove used disposable cover.

F. Complications

- Inaccurate reading (10,11,12 and 13)
- Tissue trauma
 - Rectal or colonic perforation (10,14,15)
 - Pneumoperitoneum (16)
 - Peritonitis



FIG. 6.2. Axillary temperature being taken with electronic probe thermometer. The probe is held perpendicular to the patient, and the arm is held securely against the side of the chest.

- Risk of trauma to the tympanic membrane
- Thermometer housing unit can transmit infection; disinfect after each use.

Continuous Temperature Monitoring

A. Purpose

- Provides reliable continuous monitoring of neonatal body temperature.
- Provides trend of temperature over time.
- Provides automated environmental control (Fig. 6.3).
- B. Background
 - Sites used may be surface, e.g., skin over the abdomen, or core, e.g., rectum, esophagus.
 - Thermistor probes (most widely used)
 - The thermistor is a resistive device, having a high negative temperature coefficient of resistance, so that its resistance decreases proportionately as the temperature increases.
 - As the resistance of the thermistor changes, the electrical current flowing through the probe changes proportionally.
 - The level of current detected by the electronic monitor is converted to thermal units.
 - Thermocouple probes
 - The thermocouple probe is a very small bead made up of the junction of two dissimilar metals.
 - The bead generates a very small voltage proportional to temperature.
 - The voltage generated by the bead is measured by the monitor and converted to thermal units.
 - The thermocouple and the thermistor are not interchangeable.
 - Battery-powered interface devices are available that allow the use of thermocouple probes with thermistor-compatible monitors
 - Thermocouple probes are less expensive than thermistors.



FIG. 6.3. Oxygen consumption as a function of temperature gradient between skin and environment. (From Adamsons K Jr, Gandy GM, James LS. The influence of thermal factors upon oxygen consumption of the newborn human infant. J Pediatr. 1965;66:495, with permission)

C. Indications

- Continuous temperature monitoring for whole-body cooling
- Automatic control of heater output of radiant warmer or incubator

D. Contraindications

- Rectal route in tiny infants
- E. Equipment Specifications: Hardware and Consumables
 - Continuous temperature monitoring may be a component of the bedside monitor, free-standing, or incorporated into a radiant warmer or incubator.
 - Capabilities of the neonatal temperature monitor should include:
 - Resolution to $0.1 \hat{A}^{\circ} C$
 - Temperature display in both Fahrenheit and centigrade
 - Most free-standing temperature monitors are battery-powered.
 - The monitors employ a thermistor or thermocouple.
 - Monitors using thermistors are identified as YSI 400- or YSI 700-compatible.
 - YSI 400-compatible probes are single-element devices.
 - YSI 700-compatible probes are dual-element devices.
 - YSI 400 and YSI 700 probes are physically identical and are available in the same configurations but are electrically different and will not work interchangeably.
 - Monitors using thermocouple probes are identified as such, and the probe connection is different from the thermistor type.

- Probes for both thermistors and thermocouples are available in different configurations for different sites. For example:
 - Surface skin probe
 - Tympanic membrane thermocouple probe

F. Precautions

- Do not apply skin probes to broken or bruised skin.
- Do not apply skin probes over clear plastic dressings.
- Do not use fingernails to remove skin surface probes.
- Do not force core probes during insertion.
- Do not reuse disposable probes.
- Shield skin probe with reflective pad if used with radiant warmer or heat lamp.
- When using servocontrol mechanisms for environmental control, take intermittent temperatures at other sites to monitor effectiveness (17).
- Do not use core temperature to servoregulate the patient's environment (18).

G. Technique

- Skin surface probe (Table 6.1)
 - \circ $\,$ Prepare the skin using an alcohol pad to ensure good adhesion to the skin.
 - Cover probe with a reflective cover pad (foil-covered foam adhesive pad, incorporated in the disposable probe) (Fig. 6.4). Probe must be covered with an aluminum foil disk to reflect back added heat from devices such as radiant warmers, phototherapy lights, infrared warming lights, and any other external radiant heat-generating sources (19).
 - Apply probe over the liver in the supine infant.
 - \circ $\;$ Apply probe to the flank in the prone infant.
 - \circ Ensure that skin probe is free of contact with bed (Fig. 6.5).

TABLE 6.1 Site for Temperature MonitoringSiteRange ($\hat{A}^{\circ}C$)Application

	_	
Surface		
1. Abdomen over liv	er36.0–36.5	Servocontrol
2. Axillary	36.5–37.0	Noninvasive approximation of core temperature
Core		
1. Sublingual	36.5–37.5	Quick reflection of body change
2. Esophageal	36.5–37.5	Reliable reflection of changes
3. Rectal	36.5–37.5	Slow reflection of changes

• Application of core probe (Table 6.1)

- Choose probe size according to site, i.e., rectum or esophagus.
 - Esophageal probe
 - Does not need lubrication prior to placement
 - Estimate the length of insertion, using the sum of the distance from the mouth to the tragus of the ear and the distance from the ear to the xiphoid.
 - Insert probe through nostril until the desired length is reached.



FIG. 6.4. Skin probe properly placed on infant (note that probe has protective foil cover and lies flat on the skin surface).

- Rectal probe
 - Lubricate probe before placing in rectum.
 - Probe should be placed approximately 3 cm beyond anal sphincter; avoid further advancement to avoid risk of perforation.
- Do not force either probe.
- Connect the probe to the monitor.
- Monitor energy output changes.
- Reposition or replace the probe if temperature recorded does not correlate with that recorded using an electronic thermometer. Skin surface temperature will be cooler than core temperature.



FIG. 6.5. Newborn infant with skin probe free of contact from bed surface. H. Complications

- Tissue trauma caused by core temperature probe
 - Rectal or colonic perforation
 - Pneumoperitoneum
 - Peritonitis
- Unsafe environmental temperature control caused by unshielded skin probes or loosely adhered probe, when monitoring is used to servoregulate temperature (Table 6.2).

TABLE 6.2 Potential Pitfalls of Servocontrolled Heating Devices

	Skin << Core	Skin > Core	Skin > Core
Increased heater output	Cold stress	Dislodged probe (early)	Dislodged probe
	Shock	Servo fails to shut off	(late)
	(vasoconstricted)	Vasodilators (e.g.,	Servo fails (late)
	Hypoxia	tolazoline)	
	Acidosis	Shock (vasodilated)	
Decreased heater output			
Probe uninsulated (radiant heat)			
Servocontrol malfunction			
Onset of fever			
Baby overheated			
Fever			
Internal cold stress (e.g.,			
unheated			
endotracheal oxygen, exchange			
transfusion)			

References

1. AAP Committee on Environmental Health. Mercury in the environment: implications for pediatricians. Pediatrics. 2001;107: 197.

2. Sganga A, Wallace R, Kiehl E, et al. A comparison of four methods of normal newborn temperature measurement. Am J Matern Child Nurs. 2000;25:76.

3. Craig JV, Lancaster GA, Taylor S, et al. Infrared ear thermometry compared with rectal thermometry in children: a systematic review. Lancet. 2002;360:603.

4. Siberry GK, Diener-West M, Schappell E, et al. Comparison of temple temperatures with rectal temperatures in children under two years of age. Clin Pediatr. 2002;41:405.

5. Hicks MA. A comparison of the tympanic and axillary temperatures of the preterm and term infant. J Perinatol. 1996;16:261.

6. Weir MR, Weir TE. Are "hot†ears really hot? Am J Dis Child. 1989;143:763.

7. Haddock B, Vincent P, Merrow D. Axillary and rectal temperatures of full-term neonates: are they different? Neonatal Network. 1986;5:36.

8. Stephen SB, Sexton PR. Neonatal axillary temperatures: increases in readings over time. Neonatal Network. 1987;5:25.

9. Mayfield SR, Bhatia J, Nakamura KT, et al. Temperature measurements in term and preterm neonates. J Pediatr. 1984;104:271.

 Greenbaum EI, Carson M, Kincannon WN, et al. Hazards of temperature taking. Br Med J. 1970;3:4.
 Greenbaum EI, Carson M, Kincannon WN, et al. Mercury vs. electronic thermometers. Health Devices. 1972;2:3.

12. Weiss ME, Poelter D, Gocka I. Infrared tympanic thermometry for neonatal temperature assessment. J Obstet Gynecol Neonat Nurs. 1994;23:798.

13. Ferguson GT, Gohrke C, Mansfield L. The advantages of the electronic thermometer. Hospitals. 1971;45:62.

14. Merenstein GB. Rectal perforation by thermometer. Lancet. 1970;1:1007.

15. Frank JD, Brown S. Thermometers and rectal perforations in the neonate. Arch Dis Child. 1978;53:824.

16. Greenbaum EI, Carson M, Kincannon WN, et al. Rectal thermometer-induced pneumoperitoneum in the newborn. Pediatrics. 1969;44:539.

17. Belgaumbar TK, Scott K. Effects of low humidity on small premature infants in servocontrol incubators. Biol Neonate. 1975; 26:348.

18. Friedman M, Baumgart S. Thermal regulation. In: MacDonald MG, Mullett MD, Seshia MMK, eds. Neonatology: Pathophysiology and Management of the Newborn. 6th ed. Philadelphia: Lippincott Williams & Wilkins; 2005:445.

19. Dodman N. Newborn temperature control. Neonatal Network. 1987;5:19.

7-Cardiac/Respiratory Monitoring

Rebecca J. Eick

The monitoring of vital signs in neonates provides an important indicator of overall well-being. Progress in computer technology has facilitated the development of bedside monitors that can integrate multiple monitoring parameters into a single system. This chapter covers the fundamentals of cardiac and respiratory monitoring.

Cardiac Monitoring A. Purpose

- To provide reliable and accurate monitoring of neonatal cardiac activity
 - Provide trends of heart rate over time

- Monitor beat-to-beat heart rate variability (1,2)
- To allow assessment and surveillance of critically ill neonates
- To provide early warning of potentially significant changes in heart rate by identification of heart rates above or below preset alarm limits
- To identify bradycardia (with or without associated apnea) in at-risk infants

B. Background

- Electrical activity of the heart is detected using impedance technology through skin surface electrodes (3).
- The low-level electrical signal is amplified and filtered to eliminate interference and artifacts.
- The electrical signal, defined in millivolts, is displayed as an electrocardiogram (ECG) tracing.
- R-wave detection from the QRS complex is used to calculate heart rate.
- The typical three-lead configuration (i.e., leads I, II, III) provides alternative vectors for ECG analysis.

C. Contraindications

None

D. Limitations

- The three-lead ECG is most useful for long-term continuous cardiac monitoring; more detailed cardiac evaluation (i.e., assessment of hypertrophy or axis) or the identification of abnormal cardiac rhythms may require complete 12-lead ECG with rhythm strip.
- Close proximity of electrodes in extremely small infants may cause difficulty with signal detection.

E. Equipment

Hardwareâ€"Specifications

- The monitoring system should have the appropriate frequency response and sensitivity to track the fast and narrow QRS complex of the neonate accurately.
- Heart rate is processed on a beat-to-beat basis with a short updating interval.
- Default heart rate alarm limits should be tailored to the neonatal population.
 - Low heart rate (bradycardia) limit of 100 beats/min (Note: Some term infants may have resting heart rates of 80 to 100 beats/min, requiring lower bradycardia alarm settings.)
 - \circ $\;$ High heart rate (tachycardia) limit of 175 to 200 beats/min $\;$
- Monitor displays
 - Cathode-ray tube (CRT)
 - Has highest resolution and best definition
 - Display can be either color or monochrome
 - Liquid-crystal display (LCD)
 - Flat, thin display monitor
 - Resolution may be suboptimal for fast and narrow QRS complex of neonate
 - Back-lighting is necessary for viewing in low-light environments
 - Unlike with CRT, viewing angle is critical
- Heart rate displayed as alphanumeric part of waveform display or in a separate numerical display window
- Recorder (optional)
 - Electronic memory
 - Real-time ECG
 - Delayed ECGâ€"stored retrospective display used primarily for review of short time interval prior to the occurrence of an alarm
 - Printed record of ECG trend information

- Typically used to document selected segments of ECG tracings such as periods associated with alarms or abnormal rhythms
- Monitors may have dedicated printers (often integrated into monitor cases)
- Central monitoring stations can provide remote access to information from all networked monitor units with printing capabilities.
- Units available for both bedside and transport monitoring (Figs. 7.1 and 7.2)
 - o Transport monitors typically smaller and battery-powered
 - Similar capabilities regarding parameter availability, but monitor-specific



FIG. 7.1. Typical multiparameter neonatal bedside monitor. (Courtesy of Philips Medical Systems.) Consumablesâ€"Specifications

- Disposable neonatal ECG electrodes
 - Silver–silver chloride electrodes are available in a variety of forms designed specifically for the neonatal population.
 - Patient contact surfaces of electrodes are coated in adhesive electrolyte gel, which acts as conductive medium between the patient and the metal lead while preventing direct patient contact with the metal.
 - Typical commercially available neonatal leads incorporate silver–silver chloride electrodes directly onto paper, foam, or fabric bodies with integrated lead wires.
 - Less commonly, adhesive electrode pads are separate from lead wires, which connect to the electrodes via clips.
 - ECG limb plate electrodes may be used rarely, when the application of chest leads would interfere with resuscitation or the performance of other procedures. Use of electrode gel as a conductor at the skin interface (rather than alcohol pads) is imperative in such cases.
 - Characteristics to consider in electrode selection:
 - Adherence to skin of an active infant
 - Quality of signal attained
 - Minimal skin irritation

- Ease of removal using water or adhesive remover without damage to or removal of skin
- Performance in the warm, moist environment of an infant incubator
- Adhesive–skin interaction under overhead infant warmers



FIG. 7.2. Typical multiparameter neonatal transport monitor with integrated printer. (Courtesy of Philips Medical Systems.)

- Lead wires and patient cable
 - All cables should be clean and the insulation should be free of nicks or cuts.
 - Lead wires should lock or snap into the patient cable, preventing easy disconnections.
 - If using electrodes that attach via clips, use infant/pediatric lead wires with small electrode clipsâ€"standard adult-size clips will place too much torsion on the infant electrode, tugging on the skin and possibly peeling off the electrode.

F. Precautions

- Do not leave alcohol wipes under electrodes as conductors.
- Do not apply electrodes to broken or bruised skin.
- Do not apply electrodes to clear film plastic dressingsâ€"dressing will act as an insulator between the skin and the electrode.
- To avoid skin damage, do not use fingernails to remove electrodes.
- Secure the patient cable to the patient's environment to prevent excessive traction.
- Use only monitors that have been checked for safety and performanceâ€"usually indicated by a dated sticker on the monitor.
- Do not use monitors with defects such as exposed wires, broken or dented casing, broken knobs or controls, or cracked display.
- Monitor alarms should prompt immediate patient assessment.
 - Note alarm indication (i.e., tachycardia or bradycardia).
 - Treat patient condition as necessary or correct the source of any false alarm.

• If alarm is silenced or deactivated during the course of patient evaluation, it should be reactivated prior to leaving the patient's bedside.



FIG. 7.3. Basic electrode placement and lead vectors for optimal electrocardiogram signal detection. Right arm/left arm positions also provide maximal signal for impedance pneumography. G. Techniques

- Familiarize yourself with the monitor prior to beginning.
- Electrode and lead wire placement: Although you should refer to the monitor manufacturer's placement instructions, general electrode placement guidelines follow.
 - Skin preparation: Skin should be clean and dry to provide the best electrode-to-skin interface.
 - Wipe skin with an alcohol pad and allow to dry thoroughly.
 - Avoid the use of tape to secure electrodesâ€"for optimal performance and proper electrical interface, electrodes must adhere directly to skin.
 - Basic three-lead configuration for electrode placement (for electrodes with integrated lead wires) (Fig. 7.3)
 - Right arm (white): right lateral chest at level of the nipple line
 - Left arm (black): left lateral chest at level of the nipple line
 - Left leg (red or green): left lower rib cage
 - Although this configuration allows the use of the same electrodes to monitor both ECG and respiration, optimal ECG signal may be obtained when the right arm lead is at the right mid-clavicle and the left leg lead is at the xiphoid (4).
 - If not using electrodes with integrated wires, place electrode pads in basic three-lead configuration as above, then connect lead wires via electrode clips.
 - White lead (right arm) to right chest electrode
 - Black lead (left arm) to left chest electrode
 - Red or green lead (left leg) to left lower rib cage electrode
- Turn monitor onâ€"most monitors will conduct an automatic self-test.

- Connect the patient cable to the monitor.
- Select the lead that provides the best signal and QRS size (lead II is usual default) (Fig 7.4).
 - Ensure that heart rate correlates to QRS complexes seen on displayâ€"make sure that the QRS detector is not counting high or peaked T or P waves.
- Verify that low and high heart rate alarms are set appropriately.





FIG. 7.4. Typical electrocardiogram tracings: lead I (top), lead II (middle), and lead III (bottom). H. Complications

- Skin lesions (rare)
 - o Irritation from alcoholâ€"may occur with even short-term application to immature skin
 - o Trauma caused by rubbing with excessive vigor during skin preparation
 - o Irritation from incorrectly formulated electrode gel
 - Secondary effects of skin breakdown
 - Cellulitis or abscess formation
 - Increased transepidermal water losses
 - Hypo- or hyperpigmented marks at sites of prior irritation or inflammation (Fig. 7.5)



FIG. 7.5. Residual hyperpigmented marks on the extremities present more than 1 year after application of electrocardiogram leads for cardiorespiratory monitoring. TABLE 7.1 Steps to Minimize Artifact Interference

D	T 4-	
Problem	1 reati	nent
Poor electrode	5.	Gently clean skin with alcohol wipe and allow to dry prior to
contact/connection		electrode reapplication
	6.	Check electrode/cable connections
Dried electrode	Replac	e
Equipment interference	2.	Systematically turn off one piece of adjacent equipment at a time while observing monitor for improvement in signal quality
	8.	Once source for interference identified, increase distance between that equipment and patient while rerouting power cords and cables as necessary
	9.	If above maneuver unsuccessful, replace equipment
60-hertz interference	10. 11. 12.	Follow procedure for poor electrode contact Replace patient cable If 1 and 2 unsuccessful, try alternate monitor

- Erroneous readings caused by artifacts (5) (Table 7.1)
 - Electrical interference
 - Sixty-cycle electrical interference (frequency of typical power lines)
 - Interference from other equipment used in the patient's immediate environment
 - Electrical spike may be generated when certain types of polyvinyl chloride tubing are mechanically deformed by infusion pump devicesâ€"spikes appear as ectopic beats on the monitor (rare) (6).
 - o Decreased signal amplitude with motion artifact
 - Poor electrode contact or dried electrode gel
 - Incorrect vectors because of inaccurate lead placement (Fig. 7.6)
 - Inappropriate sensitivity settings

- Monitor or cable failure
 - Hardware or software failure
 - Cable disconnection
- Alarm failure
 - False alarms (either tachycardia or bradycardia) resulting from inaccurate interpretation of heart rate
 - Inappropriate alarm parameters for patient







FIG. 7.6. Normal P-, QRS-, and T-wave detection. Top: Lead II tracing with electrodes properly placed. Note normal P-, QRS-, and T-wave detection. Middle: Lead II tracing with electrodes close together on anterior chest wall. Note altered QRS- and decreased T-wave amplitude. Bottom: Lead II tracing with electrodes placed lateral on the abdomen. Note decreased wave amplitude and flattened P wave. Respiratory Monitoring

A. Purpose

- Reliable and accurate monitoring of neonatal respiratory activity
 - Trending of respiratory activity over time
 - Detection of apnea
- Assessment and surveillance of critically ill neonates
- To provide early warning of potentially significant changes in respiratory rate by identifying respiratory rates above or below preset alarm limits

B. Background

- Measurement of transthoracic impedance is the most commonly used method for determining respiratory rate (7).
 - A low-level, high-frequency signal is passed through the patient's chest via surface electrodes.
 - Typically utilizes the same electrodes as are used for cardiac monitoring
 - Signal path usually from right arm (white) to left arm (black) electrodes, although some monitors may use right arm (white) to left leg (red or green) (Fig. 7.7)
 - \circ $\;$ Impedance to the high-frequency signal is measured.
 - Impedance is the electrical resistance to the signal.
 - Changes in lung inflation cause an alteration in the density of the chest cavity, which is detected as a change in impedance.
 - Changes in impedance modulate a proportional change in the amplitude of the high-frequency signal.
 - The change in impedance, as seen by the modulation of the high-frequency signal, is detected and quantified by the monitor and recorded as breaths per minute.
 - The monitor has an impedance threshold limit below which changes in impedance are not counted as valid respiratory activityâ€"cardiac pumping with associated changes in pulmonary blood flow will also cause changes in thoracic impedance (usually much smaller changes than those associated with respiration).



FIG. 7.7. Transthoracic impedance pneumography: diagrammatic representation of the path of the high-frequency signal between chest wall electrodes. Most monitors transmit signal right arm (white) \hat{a}^{\dagger} ' left arm (black), less commonly right arm (white) \hat{a}^{\dagger} ' left leg (red).

C. Contraindications None D. Equipment Hardwareâ€"Specifications

- Equipment is the same as that for cardiac monitoring; multiparameter monitors incorporate both cardiac and respiratory monitoring into single units.
- Respiratory monitoring parameters
 - Low-level threshold (for impedance) for breath validation should not be below 0.2 to minimize cardiogenic artifact.
 - Coincidence alarm with rejection applies when respiratory rate being detected is equal to the heart rate activity being detected by the cardiac portion of the system.
 - Default limits should be tailored to the neonatal population.
 - Adjustable apnea time-delay setting (length of apnea in seconds before alarming)
 - Typical apnea time delay is 15 to 20 seconds.

Consumablesâ€"Specifications

• Same as for cardiac monitor

E. Precautions

- Include previously discussed precautions for cardiac monitoring
- Muscular activity may be interpreted as respiration, resulting in failure to alarm during an apneic episode (see G3a following).

F. Technique

- Same as for cardiac monitor
- Ensure that the respiratory waveform correlates to the true initiation of inspiration.
- Move right and left arm electrodes up toward the axillary area if detection of respiration is poor due to shallow breathing.
- Set desired low and high respiratory rate and apnea delay alarm limits.

G. Complications

- Skin lesions (see H1 under Cardiac Monitoring)
- Monitor or cable failure
 - Hardware or software failure
 - Cable disconnection
- Alarm failure

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- False-positive "respiratory†signal in the absence of effective ventilation
 - Chest wall movement with airway obstruction (obstructive apnea)
 - Nonrespiratory muscular action (i.e., stretching, seizure, or hiccups) producing motion artifact (Fig 7.8)
- False apnea alarm despite normal respiratory activity
 - Improper sensitivity not detecting present respiratory activity
 - Incorrect electrode placement
 - Loose electrodes
 - Inappropriate alarm parameters for patient
- Accurate assessment of respiratory rate not practical when using high-frequency ventilatory modes





FIG. 7.8. Tracings of artifacts affecting electrocardiogram/ respiratory tracings. Top: Loose electrode affected by motion. Bottom: Motion artifact caused by patient's moving arm coming in contact with chest electrodes (note change in respiratory frequency signal).

Cardiorespirograph Monitoring

A. Definition

• Graphical representation of heart rate and respiratory rate over time

B. Purpose

- Monitoring of infants for identification and quantification of heart rate and respiratory activity, with detection of apnea, periodic breathing, and bradycardia
- Identification of chronologic relationships between bradycardia and apnea
- Many systems also provide continuous S_aO₂ information to allow correlation with desaturation events.

C. Background

- Heart rate is plotted graphically as beats per minute (Y axis) versus time (X axis).
- Respiratory waveform is compressed to allow display of time range.
- Short-term trending allows constant updating as the oldest information is displaced (typically based on a 2-minute window of time).
- Time relationship between heart rate and respiratory activity is maintained.
 - Allows for visualization of entire apneic episodes and identification of precipitating factors (e.g., a drop in respiratory rate may precede bradycardia)

• Inclusion of S_aO_2 allows identification of temporal relationship for desaturation events (S_aO_2 is plotted in the same fashion as the heart rate on a second Y axis)

D. ContraindicationsNoneE. EquipmentStandard features of most neonatal monitors

References

1. Cabal LA, Siassi B, Zanini B, et al. Factors affecting heart rate variability in preterm infants. Pediatrics. 1980;65:50.

2. Valimaki IA, Rautaharju PM, Roy SB, et al. Heart rate patterns in healthy term and premature infants and in respiratory distress syndrome. Eur J Cardiol. 1974;1:411.

3. Di Fiore JM. Neonatal cardiorespiratory monitoring techniques. Semin Neonatol. 2004;9:195.

4. Baird TM, Goydos JM, Neuman MR. Optimal electrode location for monitoring the ECG and breathing in neonates. Pediatr Pulmonol. 1992;12:247.

5. Jacobs MK. Sources of measurement error in noninvasive electronic instrumentation. Nurs Clin North Am. 1978;13:573.

6. Sahn DJ, Vaucher YE. Electrical current leakage transmitted to an infant via an IV controller: an unusual ECG artifact. J Pediatr. 1976;89:301.

7. Hintz SR, Wong RJ, Stevenson DK. Biomedical engineering aspects of neonatal monitoring. In: Martin RJ, Fanaroff AA, Walsh MC, eds. Fanaroff and Martin's Neonatal-Perinatal Medicine: Diseases of the Fetus and Infant. 8th ed. Philadelphia: Mosby; 2006:609.

8-Blood Pressure Monitoring

M. Kabir Abubakar

Accurate monitoring of blood pressure is essential for the optimal management of ill infants. Blood pressure is one of the most frequently measured physiologic parameters to evaluate clinical stability in critical care. The recognition and treatment of abnormal blood pressure states has significant prognostic implications in neonatal intensive care (1).

Noninvasive (Indirect) Methods

There are two types of noninvasive blood pressure measurements: auscultatory measurement (manual noninvasive) and oscillatory arterial blood pressure measurement (automatic noninvasive). Auscultatory Measurement (Manual Noninvasive)

Manual noninvasive auscultatory measurement employs the sphygmomanometer, which is simple, inexpensive, and allows for intermittent blood pressure measurements. A. Background

- This technique employs a blood pressure cuff, insufflator, manometer, and stethoscope.
- The sphygmomanometer uses a pneumatic cuff to encircle the upper arm or leg and a pressure gauge (manometer) to register the pressure in the cuff.
- There are two types of manometers:
 - Mercury (mercury column)
 - Aneroid (mechanical air gauge)
- The encircling pneumatic cuff is inflated to a pressure higher than the systolic pressure in the underlying artery. The cuff pressure compresses the artery and stops blood flow.

• A stethoscope placed distal to the cuff, over the occluded artery, will pick up the Korotkoff sounds as the pressure of the cuff is decreased to the point at which blood flow resumes through the artery.

Korotkoff sounds are the noise generated by blood spurting from the compressed artery, producing turbulence and vibration within the vessel.

- As the pressure in the cuff is reduced from above the systolic pressure, five characteristic Korotkoff sounds are heard:
 - Korotkoff I is a sharp thud.
 - Korotkoff II is a loud blowing sound.
 - Korotkoff III is a soft thud.
 - Korotkoff IV is a soft blowing sound.
 - Korotkoff V is silence.
- An 8- to 9-MHz Doppler device can be used in place of a stethoscope. This device will detect only systolic blood pressure levels.

B. Indications

- Measurement of blood pressure in stable infants or when invasive blood pressure measurement is not required or unavailable
- When only intermittent blood pressure measurements are required

C. Contraindications

- Profound edema in the limb to be measured; will affect result
- Decreased perfusion, ischemia, or infiltrate in limb
- Peripheral venous/arterial line in limb

D. Limitations

- Provides only intermittent blood pressure measurements
- Manual measurement cumbersome in small infants
- Accuracy depends on ability to recognize Korotkoff sounds and may be user-dependent
- Pressure may not be detectable in low perfusion state or shock. Do not assume that it is simply an equipment problem; use clinical correlation.
- Pressure is not detectable or is inaccurate in neonates experiencing convulsions or tremors.
- Measures only systolic and diastolic blood pressure; mean blood pressure measurement not available
- Can be used only to measure pressure in the upper arm or thigh
- Inaccurate measurements (Table 8.1)
 - Defective manometer
 - \circ $\;$ Air leaks at the tubing connections or from dry, rotted tubing
 - Inappropriate cuff size
 - Cuff applied loosely
 - Rapid deflation of cuff
 - Active or agitated patient

TADLE 0.1 Sources of Error in munect blood rressure measuren	asuremen	Me	Pressure	d	Bloo	ndirect	in	Error	of	Sources	3.1	E 8	BL	٢A
--	----------	----	----------	---	------	---------	----	-------	----	---------	------------	-----	----	----

Problem	Effect on Blood Pressure	Precaution
Defective manometer	Falsely low values	1. Check level of mercury at zero cuff
1. Air leaks		pressure
2. Improper valve funct	tion	2. Check for cleared definition of meniscus
3. Dry, rotted tubing		3. Verify that pressure holds when tightened
4. Loss of mercury		
Inappropriate cuff size		Verify appropriately sized cuff
1. Too narrow	1. Falsely high values	
2. Too wide	2. Falsely low values	
Cuff applied loosely	Falsely high values owing to ballooning of bag and narrowing of effective surface	Apply cuff snugly
Cuff applied too tightly	Inaccurate reading owing to impedance of flow through artery	Apply cuff snugly without undue pressure
Rapid deflation of cuff	 Falsely low values owing to inaccurate detection of beginning of sounds or Falsely high values owing to inadequate equilibration between cuffpressure and manometer pressure 	Deflate cuff at rate of 2–3 mm Hg/s
Active or agitated patient	Variable	Recheck when patient is quiet

TABLE 8.2 Neonatal Cuff

Cuff No. (Size)	Limb Circumference (cm)	
1	3–6	
2	4–8	
3	6–11	
4	7–13	
5	8–15	

From American Academy of Pediatrics Task Force Pressure Control: Report. Pediatrics. 1977;59:797, with permission.

E. Equipment

- Neonatal cuff (Table 8.2). Select a cuff that will fit comfortably around the upper arm or thigh; the inflatable bladder should completely encircle the extremity without overlapping. The width should be 90% of the limb circumference at the midpoint (2).
- Mercury manometer or aneroid-type gauge
- Appropriate size stethoscope with diaphragm or Doppler system

F. Precautions (Table 8.1)

- Carefully select the appropriate cuff size, because incorrect size can significantly alter the blood pressure recorded (3).
 - Cuff too small, blood pressure will be higher; cuff too large, blood pressure will be lower
- Check functional integrity of manometer
- Check integrity of cuff for leaks
- Check speed of cuff deflation: If deflation is too rapid, accuracy may be compromised.
- Patient must be quiet and still during measurements.
- For optimal infection control, use disposable cuff issued to the patient.

G. Technique

- Place the infant supine, with the limb fully extended level with the heart.
 - Measure the circumference and select the appropriate size cuff for the limb.
 - Neonatal cuffs are marked with the size range (Fig. 8.1).
 - When the cuff is wrapped around the limb, the end of the cuff should line up with the range mark.
 - \circ $\,$ If the end of the cuff falls short of the range mark, the cuff size is too small.
 - \circ If the end of the cuff falls beyond the range mark, the cuff size is too large.
- Apply the cuff snugly to the bare limb, above the elbow or knee joint.
- Place the stethoscope or Doppler over the brachial artery for the upper arm or above the popliteal artery for the thigh.
- Inflate the cuff rapidly to a pressure 15 mm Hg above the point at which the brachial pulse disappears.
- Deflate the cuff slowly.
- The pressure at which a sound is first heard is the systolic pressure (Korotkoff I). The pressure at which silence begins corresponds to the diastolic pressure (Korotkoff V). The pressure should be measured to the nearest 2 mm Hg.

In patients in whom the sounds do not disappear, the point at which the sounds change abruptly to a muffled tone can be accepted as an approximation of the diastolic pressure but will be slightly higher than true diastolic pressure.


FIG. 8.1. Properly sized infant blood pressure cuff. Arrow indicates cuff range mark. H. Complications

- Perfusion in the limb may be compromised if the cuff is not completely deflated.
- Nosocomial infection may result from using the same cuff for more than one patient.
- Prolonged or repeated cuff inflation has been associated with ischemia, purpura, and/or neuropathy.
- Cuff inflation will interfere with pulse oximetry measurement in the same limb.

Oscillometric Measurement of Arterial Blood Pressure (Automatic Noninvasive) A. Background

The oscillometric technique offers a means for measuring all arterial blood pressure parameters (systolic, diastolic, mean, heart rate) (4,5,6,7,8 and 9). This technique is referred to as NIBP (noninvasive blood pressure).

- This technique employs a blood pressure cuff interfaced to a computerized blood pressure monitor.
- The pneumatic cuff is used in the same fashion as with the auscultatory technique.
- The monitor employs a miniature computer-controlled air pump and a bleed valve to control inflation and deflation of the cuff.
- A pressure transducer interfaced to the cuff tubing senses the pressure pulsations transmitted to the cuff by the underlying artery and also the inflation pressure of the cuff.
- The system will inflate the cuff to a level above the point at which no pulsations are detected.
- As the cuff is being deflated to the level of the systolic pressure, arterial pulses are transmitted to the cuff.

The systolic pressure is assigned the value of the cuff pressure at the time pulsations were initially detected.

- With most systems, the mean pressure value is determined by the highest pulsation level detected at the lowest cuff pressure.
- The diastolic value is determined by the lowest cuff pressure before baseline arterial pulsations are detected (Fig. 8.2).
- Heart rate values are calculated by computing the mean value of the time interval between pulsations.
- Higher detection sensitivity allows this technique to be used on parts of the extremities where auscultatory methods are not possible (i.e., distal arm and lower leg).



FIG. 8.2. Determination sequence for oscillometric measurement. B. Indications

- Measurement of blood pressure in stable infants or when invasive blood pressure measurement is not required or unavailable
- When only intermittent blood pressure measurements are required

C. Contraindications

- Profound edema in the limb to be measured; will affect result
- Decreased perfusion, ischemia, or infiltrate in limb
- Peripheral venous/arterial line in place in limb

D. Limitations

- Provides only intermittent blood pressure measurements
- Pressure may not be detectable in low perfusion state or shock. Do not assume that it is simply an equipment problem; use clinical correlation.
- Pressure is not detectable or may be inaccurate in neonates experiencing convulsions or tremors.
- Inaccurate measurements (Table 8.1)
 - \circ $\;$ Air leaks at the tubing connections or from dry, rotted tubing
 - Aborted determination cycle due to air leaks in the cuff, air hoses, or connection points
 - Inappropriate cuff size

- Cuff applied loosely
- Active or agitated patient

E. Equipment

- Neonatal noninvasive blood pressure monitorâ€"display should include systolic, diastolic, mean, and heart rate values (Fig. 8.3)
- Neonatal cuff (designed for use with the specific monitor)â€"cuff may be single-tube or double-tube type, provided the appropriate adapter is used. Neonatal cuff sizes range from 1 to 5 (Table 8.2).



FIG. 8.3. Oscillometric blood pressure monitor. (Courtesy of GE Healthcare.)

F. Precautions

- Carefully select appropriate cuff size, because incorrect size can significantly alter the blood pressure value obtained.
 - Oversized cuff will yield lower blood pressure values; undersized cuff will produce higher pressure values.
- Patient must be still during measurements.
- For optimal infection control, cuffs should be for single-patient use.
- Caution should be exercised when used for preterm, very low-birthweight infants in a hypotensive state (10).

G. Technique

- Become familiar with the monitor and the equipment to be used. Be aware of the normal blood pressure changes with gestational and postnatal age (11).
- Measure the circumference of the extremity where the cuff is to be applied. Select the appropriately sized cuff for the limb (Fig. 8.4).
- Apply the cuff snugly to the limb. The cuff can be applied over a thin layer of clothing if necessary; however, a bare limb is recommended.

- Attach the monitor air hoses to the cuff. The limb from which pressure is to be measured should be level with the heart.
- Turn the monitor on and ensure that it passes the power-on self-test before proceeding.
- Press the appropriate button to start a blood pressure determination cycle.
- If the values obtained from the initial cycle are questionable, repeat step 6.

Multiple readings with similar values yield the optimal assurance of accuracy.

- If, after repeating the cycle, readings are still questionable, reposition the cuff and repeat the measurement.
- Periodic inspection of the cuff and extremity is critical to avoid problems such as cuff detachment or shift in extremity position.
- Most NIBP systems can be programmed by the user to measure blood pressure automatically at userdetermined intervals.

The interval between measurements should be as long as possible to ensure adequate circulation and minimize trauma to the limb and skin distal to the cuff.

• In infants with suspected congenital heart disease, blood pressure should be measured in all four extremities or at least the right arm and right leg for pre- and postductal comparison.



FIG. 8.4. Cuff of correct size applied to upper arm. H. Complications

- Perfusion in the limb may be compromised if the cuff is not completely deflated.
- Nosocomial infection may arise from using the same cuff for more than one patient.
- Repeated continuous cycling may cause ischemia, purpura, and/or neuropathy.
- Cuff inflation will interfere with pulse oximetry measurement and IV infusion in the same limb.

Continuous Blood Pressure Monitoring (Invasive)

A. Purpose

Accurate blood pressure measurements are essential to guide the level of circulatory support provided to critically ill infants. Direct reading through an intravascular catheter provides the most accurate measurement and is the "gold standard†in measuring blood pressure. B. Background

- Blood pressure measurement is obtained from the vascular system via a catheter that has been introduced into a vein or artery. The catheter is coupled to a pressure transducer outside the body by a fluid-filled conduit.
- Because fluid is noncompressible, it is the medium used to transmit the pressure from the vessel to the transducer.
- The tubing used to couple the catheter to the transducer is a special high-pressure tubing with low compliance that is capable of withstanding high pressures without breakage and with very little stretching.
- A blood pressure transducer is a device that converts mechanical forces (pressure) to electrical signals. There are two major types of transducers:
 - Strain gauge pressure transducer: composed of metal strands or foil that is either stretched or released by the applied pressure on the diaphragm

Applied pressure causes a proportional and linear change in electrical resistance. Problems associated with strain gauges include drift due to temperature changes (departure from the real signal value), fragility, and cost.

• Solid-state pressure transducer (semiconductor): composed of a silicone chip that undergoes electrical resistance changes because of the applied pressure.

This technology offers a superior alternative to the strain gauge.

- Because of its low cost, it has made possible inexpensive, accurate, disposable transducers.
- Because of the miniature integration on the silicone chip, the circuitry necessary to minimize temperature drift is incorporated in the device.
- Pressure transmitted to the transducer causes a micromechanical displacement of the diaphragm (receptor of mechanical energy linked to the transducer element) or directly to the transducer.
- This mechanical displacement causes a proportional change in the electrical resistance of the transducer.
- Excitation voltage applied to the transducer is responsible for the signal produced in relation to the pressure applied.
- The standard medical blood pressure transducer output rating is 5 mV/V/cm Hg.
- Miniature transducer-tipped catheters are available that do not depend on fluid-filled lines for the transmission of pressure.

Microtransducer catheters in general have better fidelity characteristics, but at a much higher cost than conventional fluid-filled systems.

• The pressure monitor processes the electrical signal generated by the transducer and converts it to blood pressure units in either millimeters of mercury or kilopascals. Part of this processing involves the detection of systolic, diastolic, and mean values.

C. Indications

- To monitor intravascular pressure continuously
 - In very small or unstable infants, particularly those with severe hypotension (shock)
 - During major procedures that could cause or exacerbate intravascular instability
 - To monitor infants on aggressive ventilator support or extracorporeal membrane oxygenation

D. Contraindications

None absolute, except for those specific to catheter placement

- E. Limitations
 - The pulse pressure waveform measured in the periphery is narrower and taller than that in the proximal aorta. Thus, systolic blood pressure in the peripheral arteries can be higher than that in proximal aorta. This amplification is greater in patients with increased vascular tone or on inotropic therapy.
 - Very–small-diameter catheters may result in underreading of systolic blood pressure.

F. Equipment

- Neonatal physiologic monitor (multiparameter monitoring system)
 - Minimum configuration should have the capability of displaying systolic, diastolic, and mean pressures.
 - It should have provision for high and low alarm settings.
- Mechanical infusion device (infusion pump)

Pressurized intravenous bag should never be used.

- Fluid to be infused via the pressure line
- Appropriately sized catheter and supplies
- Pressure-monitoring kit with integrated disposable transducer and continuous flush device (Fig. 8.5)
 - Pressure-monitoring tubing should not exceed 48 inches from the transducer to the patient connection.
 - The distal end of the kit should have a stopcock or an arterial extension no longer than 12 inches that can be used for drawing blood samples.
 - Some disposable pressure-monitoring kits offer closed-loop systems for sampling.
 - The system employs a mechanism for aspirating and holding a fixed amount of blood in the pressure tubing rather than in a syringe.
 - The distal end is equipped with a small chamber with a rubber septum that allows a self-guiding short blunt syringe adapter to penetrate and aspirate blood for the sample.

The initial volume pulled back is sufficient to ensure that the blood drawn into the sample chamber is greater than the catheter/distal tubing volume and is not diluted by the fluid being infused. The absence of stopcocks at the distal end eliminates a possible site for contamination. In addition, the blood pulled back is conserved, and the amount of fluid used to flush the sample line is reduced.



FIG. 8.5. Representative disposable blood pressure transducer setup. (1) Pressure transducer; (2) integral continuous flush device; (3) infusion port (connects to infusion pump); (4) high-pressure tubing.

G. Technique

For catheter placement, see Section 5 of the atlas, "Vascular Accessâ€



FIG. 8.6. Arterial pressure waveforms: normal arterial waveform (top); dampened arterial waveform

(middle); arterial waveform with spike caused by catheter whip or inappropriate tubing (bottom). (Note that figure demonstrates waveform appearance only and not actual pressure values.)

- Familiarize yourself with the bedside monitor and the pressure zero/calibration procedure.
- If using discrete components, assemble the pressure-monitoring circuit, maintaining the sterile integrity.
 - A basic circuit configuration will consist of a transducer dome, flush device, stopcock, pressure tubing, and an optional arterial extension set (short length of pressure tubing, less than 12 in in length, inserted between the catheter and the pressure tubing).
 - If using a pressure-monitoring kit, ensure that all the Luer-lock connections are tight and free of any defects.
 - If possible, avoid the use of intravenous tubing components in the pressure-monitoring circuit.
- Set up the infusion pump that will be used for the continuous infusion through the flush device. Continuous flush devices limit flow rates to 3 or 30 mL/h, depending on the model (12). For neonatal arterial lines, the infusion pump supplying the flush device should be set to 0.5 to 3 mL/h and should never exceed the flow rating of the flush device. When pump flow exceeds the flush device rating, it will cause an occlusion alarm in most intravenous pumps. A pump flow rate of 1 mL/h is recommended for most arterial lines.
- For circuit priming, use the solution that will be used for the continuous infusion. Prime the circuit slowly, in order to avoid trapping air bubbles in the flush device inlet. Ensure that the entire circuit and all the ports are filled and bubble-free.
- If using disposable transducers, connect the reusable interface cable to the transducer and to the monitor. Turn the monitor on.
- Secure the transducer at the patient's reference level, defined as the midaxillary line (heart level). If using transducer holders, level the reference mark on the holder at the patient's reference level.
- Connect the distal end of the circuit to the patient's catheter, ensuring that the catheter hub is filled with fluid and is bubble-free.
- Start the infusion pump. The pump rate cannot exceed the flow rate of the flush device.
- Open the stopcock connected to the transducer to air (shut off to the patient, open to atmosphere).
- Zero/calibrate the monitor according to the manufacturer's instructions.
- Close the stopcock connected to the transducer (open to the patient).
- Set the monitor pressure waveform scale to one that accommodates the entire pressure wave.
- Observe the waveform obtained. If the wave appears to be dampened (flattened, poorly defined, with slow rise time), check the circuit for air bubbles starting at the distal end. If no air bubbles are detected, then gently flush the catheter (Fig. 8.6). (For central venous waveforms, see Chapter 29.)
- Once a stable pressure reading is obtained, set the alarm limits. Mean pressure value is optimally used to set alarm limits (Fig. 8.7).
- Zero the transducer every shift or every 8 hours.
- When blood samples are drawn from the line, flushing should be done gently with a syringe using a minimal amount of heparinized saline solution.



FIG. 8.7. Pressures obtained by direct measurement through umbilical artery catheter in healthy newborn infants during first 12 hours of life. Broken lines represent linear regressions; solid lines represent 95% confidence limits. A: Systolic pressure (top) and diastolic pressure (bottom). B: Mean aortic pressure (top) and pulse pressure (systolic–diastolic pressure amplitude) (bottom). (From Versmold HT, Kitterman JA, Phibbs RH, et al. Aortic blood pressure during the first 12 hours of life in infants with birth weight 610 to 4,220 grams. Pediatrics. 1981;67:611, with permission of American Academy of Pediatrics.)

H. Complications (Table 8.3)

- Defective transducer
- Cracked Luer-lock connections, causing leaks, low pressure readings, or blood to back up in the line
- Air in the line
- Malfunctioning infusion pump not providing continuous flush, causing the line to clot off
- Defective reusable transducer interface cable (disposable transducer system)
- Erroneous readings caused by the transducer not being properly set at the patient's reference level or if it slips from the preset level

Lower readings occur when the transducer is high; higher readings occur when the transducer is low.

- Problems associated with catheters
- Tip of the catheter lodging against the wall of the vessel (will cause the pressure wave to flatten and the pressure to rise slowly as a result of the continuous infusion)
- Transducer not zeroed to atmosphere (static pressure trapped by stopcock valve and a syringe stuck in the port that should be opened to air)

Will cause lower or negative pressure readings

- Loss of blood if stopcock is left opened to the patient and air
- Fluid overload if a pressurized intravenous bag is used instead of an infusion pump and the fast flush mode is used to clear the line (11)

Problem	Cause	Prevention	Treatment
Damped pressure	Catheter tip against vessel	Usually unavoidable	Reposition catheter while
tracing	wall		observing waveform
	Partial occlusion of tip by	Use continuous infusion of 5%	Aspirate clot with syringe
	clot	dextrose in water or add	and flush with heparinized
		heparin (1 U/mL IV fluid)	saline
	Clotting in stopcock or	Flush catheter carefully after	Flush stopcock and
	transducer, or blood in	blood withdrawal and	transducer; if no
	system	reestablish IV drip; back-flush	improvement, change
		stopcocks to remove blood	components
Abnormally high or low readings	Change in transducer level	Maintain patient in same	Recheck patient and
		position for serial pressure	transducer positions
		measurements	
	Leaks in transducer system	Assemble transducer carefully,	Check all fittings, transducer
		ensuring that dome is attached	dome, and stopcock
		snugly; use Luer-lock fittings	connections
		and disposable stopcocks	T
	External vascular	Secure catheter firmly without	Loosen tape, securing
	compression	on extremity	catheter in place
	Strained transducer	Attention to stopcocks when aspirating to module	Replace transducer
	Loose dome on transducer	Attach transducer dome	Vent system and retighten
	producing low blood	securely to module	dome
	pressure without leak or		
	change in waveform ^a		
	High intrathoracic pressure	Be aware of problem	Take pressure readings off
	secondary to ventilation;		ventilation or in expiratory
	particularly influences		phase
	central venous pressure		
Damped pressure	Air bubbles in transducer	Flush transducer and tubing	Check system, rapid flush,
Without	connector tubing	carefully when setting up	attach syringe to transducer,
fluching		system and attaching to	and aspirate bubble
nusning		carefully	
No pressure	Transducer not open to	Follow routing systematic	Chack system ² f"stoncocks
available	catheter or settings on	steps for setting up system and	monitor and amplifier setup
available	monitor amplifiers	nressure measurements	monitor, and ampinier setup
	incorrectâ€"still on zero	pressure measurements	
	cal, or off		

 TABLE 8.3 Trouble-shooting for Intravascular Pressure Monitoring

^aData from Weindling AM. Blood pressure monitoring in the newborn. Arch Dis Child. 1989;64:444.

References

1. Nuntnarumit P, Yang W, Bada-Ellzey HS. Blood pressure measurements in the newborn. Clin Perinatol. 1999;26(4): 981.

2. Bickley LS, Hoekelman RA. In: Bickley LS, ed. Bates' Guide to Physical Examination and History Taking. 7th ed. Philadelphia: Lippincott Williams & Wilkins; 1999.

3. Moss AJ. Indirect methods of blood pressure measurement. Pediatr Clin North Am. 1978;25:3.

4. Kirkendall WM, Feinleib M, Freis ED, et al. Recommendation for human blood pressure determination by sphygmomanometers: Subcommittee on the American Heart Association Postgraduate Education Committee. Circulation. 1980;62:1145A.

5. Sadove MS, Schmidt G, Wu H-H, et al. Indirect blood pressure measurement in infants: a comparison of four methods in four limbs. Anesth Analg. 1973;52:682.

6. Ramsey M III. Automatic oscillometric noninvasive blood pressure: theory and practice. In: Meyer-Sabellak W, ed. Blood Pressure Measurements. Darmstadt: Steinkopff Verlag; 1990:15. P.57

7. Kimble KJ, Darnall RA Jr, Yelderman M, et al. An automated oscillometric technique for estimating mean arterial pressure in critically ill newborns. Anesthesiology. 1981; 54:423.

8. Friesen RH, Lichtor JL. Indirect measurement of blood pressure in neonates and infants utilizing an automatic noninvasive oscillometric monitor. Anesth Analg. 1981;60:742.

9. Park MK, Menard SM. Accuracy of blood pressure measurement by the Dinamap monitor in infants and children. Pediatrics. 1987;79:907.

10. Weindling AM. Blood pressure monitoring in the newborn. Arch Dis Child. 1989;64:444.

11. Lee J, Rajadurai VS, Tank KW. Blood pressure standards for very low birth weight infants during the first days of life. Arch Dis Child Fetal Neonat Ed. 1999;81:F168.

12. Morray J, Todd S. A hazard of continuous flush systems for vascular pressure monitoring in infants. Anesthesiology. 1983;38:187.

9-Continuous Blood Gas Monitoring

M. Kabir Abubakar

Pulse Oximetry

A. Definitions

- Arterial oxyhemoglobin saturation measured by an arterial blood gas analysis is referred to as SaO₂.
- Arterial oxyhemoglobin saturation measured noninvasively by pulse oximetry is referred to as SpO₂.

B. Purpose

- Noninvasive arterial oxygen saturation monitoring
- Pulse rate monitoring
- Trending of SpO₂ and pulse rate over time

C. Background

- Principles of oxygen transport
 - Approximately 98% of the oxygen in the blood is bound to hemoglobin.

The amount of oxygen carried in the blood is related directly to the amount of hemoglobin in the blood and to the partial pressure of unbound, dissolved oxygen in the blood (PO₂) (1). The relationship of arterial PO₂, in nearâ \in "term infants, to percent saturation measured by pulse oximeter is shown in Fig. 9.1.

• The relationship between blood PO₂ and the amount of oxygen bound to hemoglobin is presented graphically as an oxygen–hemoglobin affinity curve (Fig. 9.2). Percent oxygen saturation is calculated below:

Oxyhemoglobin

 $\overline{\text{Oxyhemoglobin} + \text{Deoxyhemoglobin}} \times 100$

- Principles of pulse oximetry
 - Based on the principles of spectrophotometric oximetry and plethysmography (2)
 - Determine arterial saturation and pulse rate by measuring the absorption of selected wavelengths of light

Oxygenated hemoglobin (oxyhemoglobin) and reduced hemoglobin (deoxyhemoglobin) absorb light as known functions of wavelengths. By measuring the absorption levels at different wavelengths of light, the relative percentages of these two constituents and SpO_2 are calculated.

- Utilizes a sensor composed of two light-emitting diodes (LEDs) as light sources and one photodetector as a light receiver. The photodetector is an electronic device that produces a current proportional to the incident light intensity (3).
 - One LED emits red light with an approximate wavelength of 660 nm.

Red light is absorbed selectively by deoxyhemoglobin.

• The other LED emits infrared light with an approximate wavelength of 925 nm.

Infrared light is absorbed selectively by oxyhemoglobin.

- Utilizes the different absorption of the wavelengths when transmitted through tissue, pulsatile blood, and nonpulsatile blood (Fig. 9.3).
 - The photodetector measures the level of light that passes through without being absorbed.
 - During the absence of pulse (diastole), the detector establishes baseline levels for the absorption of tissue and nonpulsatile blood.
 - With each heartbeat, a pulse of oxygenated blood flows to the sensor site.
 - Absorption during systole of both the red and the infrared light is measured to determine the percentage of oxyhemoglobin.
 - Because the measurements of the change in absorption are made during a pulse (systole), these pulses are counted and displayed as heart rate.

D. Indications

- To monitor oxygenation in infants suffering from conditions associated with:
 - o Hypoxia
 - Apnea/hypoventilation
 - Cardiorespiratory disease



FIG. 9.1. Arterial PO₂ versus pulse oximeter percent saturation in near-term newborn infants in whom pulse saturation was fixed by adjusting F_iO_2 first and then measuring P_aO_2 . Values are means $\hat{A}\pm$ SD. (From Brockway J, Hay WW Jr. Ability of pulse O₂ saturations to accurately determine blood oxygenation. Clin Res. 1988;36:227A , with permission.)

Bronchopulmonary dysplasia

Pulse oximetry is the optimal mode of monitoring for larger infants with bronchopulmonary dysplasia. Whereas PO₂ monitors may underestimate P_aO_2 in this population, oximetry has been found to be reasonably accurate for these infants (4). The pulse oximeter requires no patient preparation or calibration time, has a rapid response time, and is readily adaptable to different patient populations (5).





Hay WW Jr. Physiology of oxygenation and its relation to pulse oximetry in neonates. J Perinatol. 1987;7:309

, with permission.)



FIG. 9.3. Tissue composite showing dynamic as well as static components affecting light absorption. (From

Wukitch MW, Petterson MT, Tobler DR, et al. Pulse oximetry: analysis of theory, technology and practice. J Clin Monit. 1988;4:290

, with permission.)

- To monitor response to therapy
 - Resuscitation (6)

Pulse oximetry is also a significant adjunct to monitoring in the delivery room. With the use of pulse oximetry, SpO_2 values can be obtained within 1 minute after birth (7, 8, 9) (Fig. 9.4).

- Monitoring effectiveness of mask ventilation (10) or during placement of an endotracheal tube
- To monitor side effects of other therapy
 - \circ Suctioning
 - Positioning for laryngoscopy (6, spinal tap, etc.
- For low-birthweight infants <1,000 g (11, 12)



FIG. 9.4. Mean arterial oxygen saturation (S_aO_2) values measured by pulse oximetry from the time of cord clamping. Values are means $\hat{A}\pm$ SD. (From House JT, Schultetus RR, Gravenstein N. Continuous neonatal evaluation in the delivery room by pulse oximetry. J Clin Monit. 1987;3:96

, with permission.)

- It is optimal to use pulse oximetry for oxygen monitoring in the very low-birthweight infant because of its
- P.60
- •

noninvasiveness. Pulse oximetry can be used reliably in very low-birthweight infants with acute as well as chronic lung disease (11).

• Pulse oximetry also offers an advantage for precise fraction of inspired oxygen (F_iO_2) control during neonatal anesthesia because of the short response time to changes in SpO₂ (13).

E. Limitations

• Decreased accuracy when arterial saturation is <65%

Pulse oximetry will overestimate SpO_2 at this level; therefore, blood gas confirmation is imperative (14, 15, 16).

• Not a sensitive indicator for hyperoxemia (1)

Pulse oximeter accuracy does not allow for precise estimation of PO_2 at saturations >90%. Small changes in O_2 saturation (1% to 2%) may be associated with large changes in PO_2 (6 to 12 mm Hg) (1).

• Because pulse oximeters rely on pulsatile fluctuations in transmitted light intensity to estimate SpO₂, they are all adversely affected by movement (3, 17 and 19).

In some cases, the pulse oximeter may calculate an SpO₂ value for signals caused by movement, or it may reject the signal and not update the display. Usually, the heart rate output from the oximeter will reflect the detection of nonarterial pulsations, indicating either $\hat{a} \in \hat{c} \otimes \hat{o} \in \hat{c}$ saturation or $\hat{a} \in \hat{c} \otimes \hat{c}$ quality signal $\hat{a} \in \hat{c}$. (3). However, advances in microprocessor technology have led to improved signal processing, which makes it possible to minimize motion artifact and monitor saturation more accurately during motion or low-perfusion states.

- Significant levels of carboxyhemoglobin or methemoglobin can yield erroneous readings (carboxyhemoglobin absorbs light at the 660-nm wavelength) (18). However, carboxyhemoglobin levels of <3% will not affect the accuracy of the instrument.
- SpO₂ may be overestimated in darkly pigmented infants.

Some oximeters will give a message such as $\hat{a} \in \hat{c}$ insufficient signal detected $\hat{a} \in \hat{c}$ if a valid signal is not obtained (18, 19).

• Erroneous readings can occur in the presence of high fetal hemoglobin (18).

A smaller effect on accuracy is noted when fetal hemoglobin levels are <50% (20, 21 and 22). With a predominance of fetal hemoglobin, an SpO₂ of >92% may be associated with hyperoxemia (22). However, whereas saturations may appear adequate, PO₂ may be low enough to produce increased pulmonary vascular resistance (SpO₂/PO₂ curve shift to the left).

Because infants with chronic lung disease and prolonged oxygen dependence are older and have less fetal hemoglobin, SpO_2 readings obtained from these patients may be more accurate than those obtained from neonates with acute respiratory disorders at an earlier age (23). The same situation exists in infants who have undergone exchange transfusion because of decreased levels of fetal hemoglobin (24).

• Light sources that can affect performance include surgical lights, xenon lights, bilirubin lamps, fluorescent lights, infrared heating lamps, and direct sunlight.

Although jaundice does not account for variability in pulse oximeter accuracy (24), phototherapy can interfere with accurate monitoring. Therefore, appropriate precautions should be taken, such as covering the probe with a relatively opaque material (1).

• Do not correlate SpO₂ values with laboratory hemoximeters (18).

Most laboratory oximeters measure fractional oxygen saturation (all hemoglobin including dysfunctional hemoglobin) as opposed to functional oxygen saturation (oxyhemoglobin and deoxyhemoglobin excluding all dysfunctional hemoglobin).

• Use of normal adult values for hemoglobin, 2,3-diphosphoglycerate, and, in some cases, PCO₂ can lead to errors in the algorithm to calculate SpO₂ with some blood gas analysis instruments (18).

• Although pulse oximeters can detect hyperoxemia, it is important that type-specific alarm limits are set (25).

To avoid hyperoxemia, a minimal sensitivity of at least 95% is required.

- Pulse oximeters rely on detecting pulsatile flow in body tissues; therefore a reduction in peripheral pulsatile blood flow produced by peripheral vasoconstriction results in an inadequate signal for analysis.
- Pulse oximeters average their readings over several seconds depending on oximeter type and internal settings. Oximeters with a long averaging time may not be able to detect acute and transient changes in SpO₂.
- Venous congestion may produce venous pulsations, which can produce low readings.
- The pulse oximeter only provides information about oxygenation. It does not give any indication of the patient's carbon dioxide elimination.
- In summary, it is optimal to make some correlation between SpO₂ and PO₂ throughout a reasonable range of SpO₂ (lower, 85% to 88%; higher, 95% to 97%) before relying completely on SpO₂ for oxygen and/or respirator management (23, 26).



FIG. 9.5. Pulse oximeter. Vertical column indicates pulse. (Courtesy of Nellcor, Pleasanton, CA, USA.) F. Equipment

- Manufacturer-specific sensor and monitor (Fig. 9.5) with
 - \circ Display of SpO₂ and pulse rate and a pulse indicator
 - Adjustable alarm limits for SpO₂ as well as pulse rate
 - Battery-powered operation
- Neonatal sensor, either disposable or reusable
 - Disposable sensors have become the standard from the standpoint of infection as well as quality control.
 - Disposable neonatal sensors are available in different sizes, depending on the site to be used.

G. Precautions

• Use only with detectable pulse.

Cardiopulmonary bypass with nonpulsating flow, inflated blood pressure cuff proximal to the sensor, tense peripheral edema, hypothermia, low perfusion state secondary to shock or severe hypovolemia, and significant peripheral vasoconstriction may interfere with obtaining accurate readings (5).

- Assess the sensor site every 8 hours to be certain that the adherent bandage is not constricting the site and that the skin is intact.
- Whenever possible, the SpO₂ sensor should not be on the same extremity as the blood pressure cuff.

When the cuff is inflated, the SpO_2 sensor will not detect a pulse, will not update SpO_2 values, and will alarm. Use of ace bandages on the extremities to increase central venous return may also interfere with the function of the sensor.

- Malpositioned sensor: When a probe is not placed symmetrically, it can allow some light from the LED emitters to reach the photodetector in the sensor without going through the tissue at the monitoring site and will therefore produce falsely low readings. This is called the penumbra effect.
- To avoid possible transfer of infection, do not share pulse oximetry probes between patients.

H. Technique

- Familiarize yourself with the system before proceeding.
- Select an appropriate sensor and apply it to the patient.
 - Ear lobe, finger, toe, lateral side of the foot, or across the palm of the hand. (Placing the sensor in a position matching that of the peripheral arterial line, if present, may avoid discrepancies caused by intracardiac or ductal shunts when trying to correlate SpO₂ with arterial PO₂.)
 - For neonates 500 g to 3 kg, anterior-lateral aspect of a foot (Fig. 9.6) (1).
 - \circ For infants weighing >3 kg, use the palm, thumb, great toe, or index finger (1).
 - Align the LEDs (light source) and the detector so they are directly opposite each other.
 - Reusable sensors should be applied with nonadhesive elastic wrap.
 - Tighten sensor snugly to the skin but not so as to impede circulation. The probe should then be left in place for several seconds until extremity movement stops and the signal is stable.
 - Secure the sensor to the site to prevent tugging or movement of the sensor independent of the body part.
 - Cover the sensor to reduce the effect of intense light levels, direct sunlight, or phototherapy.
- Attach the sensor to the system interconnecting cable and turn on the monitor. (Attaching the sensor to the baby before connecting the cable to the monitor will shorten the time taken for data acquisition and display of SpO_2 information.)
- Calibration of the system is not required (internal autocalibration).
- After a short interval, if all connections are correct, the monitor will display the pulse detected by the sensor. If the pulse level is adequate, it will display

 SpO_2 and pulse rate. If the pulse indicator is not synchronous with the patient's pulse rate, reposition the probe. If, after repositioning the sensor, the pulse detector is still not indicating properly, change the sensor site.

• Once reliable operation is achieved, set the high and low alarm limits.

- Although pulse oximeters can detect hyperoxemia, it is important that type-specific alarm limits are set and a low specificity is accepted (25). Type-specific alarm limits should be guided by normative values for SpO₂ depending on postnatal age (Fig. 9.7).
- The optimal alarm limit, defined as having a sensitivity of 95% or more, associated with maximal specificity, will differ depending on which particular monitor is used.

In general, low-limit SpO₂ values are defined as 87% to 89%; higher-limit SpO₂ values are defined as 94% to 95%.

• Default (starting point) alarm limits for a newborn (26): high limit >94% to 95%; low limit <87% to 90%.

Note that SpO_2 is a more sensitive indicator of hypoxemia and decreased tissue oxygenation than is P_aO_2 . Lower alarm limits should be individualized to alert the user when the oxygenation requirements of the given patient are not satisfied.



FIG. 9.6. Disposable sensor applied to foot.



FIG. 9.7 The oxygen dissociation curve at various postnatal ages. (From Oski FA, Delivoria-Papadopolous POM. The red cell 2,3-diphosphoglycerate and tissue oxygen release. J Pediatr. 1970;77:941 , with permission.)

I. Complications

- Management based on erroneous readings caused by a misapplied sensor or conditions affecting instrument performance (27, 29)
- Burn resulting from electrical short (28)
- Limb ischemia if applied too tightly, particularly in an edematous limb

Transcutaneous Blood Gas Monitoring

Transcutaneous measurements of oxygen and carbon dioxide are useful in the neonatal intensive care unit because they provide continuous and relatively noninvasive estimation of arterial blood gases. A. Definitions

- Transcutaneous measurement of oxygen is referred to as PtcO2.
- Transcutaneous measurement of carbon dioxide is referred to as PtcCO₂.

B. Purpose

- Noninvasive blood gas monitoring of PO₂ and PCO₂
- Trending of PO₂ and PCO₂ over time

C. Background

- Transcutaneous monitoring measures skin-surface PO_2 and PCO_2 to provide estimates of arterial partial pressure of oxygen and carbon dioxide. The devices increase tissue perfusion by heating the skin and then electrochemically measuring the partial pressure of oxygen and carbon dioxide.
- Accomplished by two electrodes contained in a heated block that maintains the electrodes and the skin directly beneath it at a constant temperature (30) (Fig. 9.8)
 - Arterialized capillary oxygen levels are more accurately measured by heating the skin to establish hyperemia directly beneath the sensor.
 - The electrodes are covered with an electrolyte solution and sealed with a semipermeable plastic membrane.
- A modified Clark electrode is used to measure oxygen.
 - \circ It produces an electrical current that is proportional to PO₂.
 - \circ $\,$ Measured current is converted to PO_2 and then corrected for temperature.



FIG. 9.8. A: Principle of cutaneous PO₂ measurement by heated oxygen sensor. B: Temperature profile of cutaneous tissue. C: Cross section of cutaneous oxygen sensor. (Courtesy of Kontron Medical Instruments, Ergolding, Germany.)

- A Severinghaus electrode is used to measure CO₂.
 - pH-sensitive glass electrode
 - \circ CO₂ diffuses from the skin surface through the membrane. The CO₂ changes the pH of the electrolyte solution bathing the electrode.
 - The measured pH is converted to PCO₂ and then corrected for temperature.

Conversion of electric current and pH to PO_2 and PCO_2 , respectively, is based on conversion equations adjusted by a two-point calibration. This is part of the setup and calibration process.

D. Indications

- To approximate arterial P_aO_2 and P_aCO_2 for respiratory management
 - To monitor the effect of therapeutic ventilatory maneuvers (30, 31, 32, 33 and 34), particularly in infants who have combined oxygenation and ventilation problems
 - For stabilization and monitoring during transport (31, 35, 36)
- To reduce the frequency of arterial blood gas analysis (37)
- To determine by a noninvasive and continuous method the regional arterial oxygen tension (38, 39 and 40)
- To infer regional arterial blood flow (40, 41, 42 and 43), e.g., in the lower limbs of infants with ductdependent coarctation of the aorta

E. Contraindications

- Skin disorders (e.g., epidermolysis bullosa, staphylococcal scalded skin syndrome)
- Relative contraindications
 - The very low-birthweight infant (44, 45)
 - Severe acidosis
 - Significant anemia
 - Decreased peripheral perfusion
 - $\circ \quad P_{tc}O_2 \text{ may underestimate } P_aO_2 \ (46).$





F. Equipmentâ€"Specifications

- Transcutaneous monitor components
 - Dual electrode
 - Electrode cleaning kit
 - Electrolyte and membrane kit
 - Contact solution

- Double-sided adhesive rings
- Calibration gas cylinders with delivery apparatus
- Digital display shows values for $P_{tc}O_2$, $P_{tc}CO_2$, and site of sensor (Fig. 9.9).
- Monitor with controls for both high and low alarm limits, and for electrode temperature. The monitor may also have a site placement timer that will alarm as an indication to change the site of the electrode.

G. Precautions

- Be aware that:
 - Equilibration requires approximately 20 minutes once the electrode is placed, with the response time for $P_{tc}O_2$ being much faster than that for $P_{tc}CO_2$. Therefore, management changes based on transcutaneous values should be guided by values that have been consistent for at least 5 minutes.
 - \circ Periodic correlation with PO₂ from appropriate arterial sites is recommended (46).
 - $P_{tc}O_2$ may underestimate P_aO_2 in the infant with hyperoxemia ($P_aO_2 > 100 \text{ mm Hg}$), with reliability of $P_{tc}O_2$ measurement decreasing as P_aO_2 increases (47, 48, 49 and 50).
 - \circ P_{tc}O₂ may underestimate P_aO₂ in older infants with bronchopulmonary dysplasia (51, 52).
 - Transcutaneous blood gas measurements are affected by the state of the infant.

Values during feeding or active sleep can be lower (53).

- \circ Pressure on the sensor (e.g., infant lying on sensor) may restrict blood supply, resulting in falsely low P_{tc}O₂ values.
- Manufacturers' parts are not interchangeable. Only supplies of the same brand and designated for the monitor should be used.
- To avoid skin burns, change electrode location at least every 4 hours.
- $P_{tc}O_2$ may underestimate P_aO_2 in the presence of:
 - Severe acidosis
 - Severe anemia
 - Decreased peripheral perfusion

H. Technique

- Familiarize yourself with the system before proceeding.
 - Perform routine electrode maintenance, if there is any question as to the status of the electrode.
 - Remove the membrane, rinse the electrode with deionized water, and dry with a soft lint-free tissue or gauze.
 - Clean the electrode using the solution provided in the cleaning kit; abrasive compounds or materials should never be used (they will permanently damage the electrode).
 - \circ $\;$ Rinse the electrode with deionized water and dry with lint-free tissue.
 - Apply the electrolyte solution.
 - Place a new membrane on the electrode. Avoid finger contact, and always handle the membrane inside its protective package or with plastic tweezers.
- Perform two-point gas calibration using the device-specific apparatus, as per manufacturer's instruction.
- Use an alcohol pad to clean and degrease the skin site where the sensor is to be placed.
- Apply double-sided adhesive ring to the sensor.
- Apply one drop of contact solution to the skin site.
- Peel the protective backing from the adhesive ring, place the sensor on the skin over the contact solution, and press the sensor to the skin.
 - \circ $\;$ For best results, place the sensor on a location with good blood flow.

- Appropriate sites include the lateral abdomen, anterior or lateral chest, volar aspect of the forearm, inner upper arm, inner thigh, or posterior chest (Fig. 9.10) (54).
- Although large differences between pre- and postductal P_aO₂ values are uncommon in premature infants with hyaline membrane disease, preductal location of the electrode is optimal for prevention of hyperoxemia (55).
- Choose a site devoid of hair.
- Avoid bony prominences.
- Avoid areas with large surface blood vessels (Fig. 9.10).



FIG. 9.10. Cutaneous PO₂/PCO₂ sensor applied to the back.

- Secure the sensor cable to prevent tugging of the electrode when the cable is manipulated.
- Allow 15 to 20 minutes for site equilibration before taking readings.
- Note the time at which the sensor was placed on the skin, so that the site can be changed after a 4-hour period (maximum site time). When changing the sensor site:
 - Use an alcohol pad to help loosen the adhesive and peel gently from the skin.
 - Inspect the skin site for signs of sensitivity to heat or to the adhesive. In the event of skin irritation, either lower the sensor temperature or change the site more frequently; mild erythema after sensor removal is typical.
 - Peel adhesive ring off the sensor.
 - Flush the membrane surface with de-ionized water.
 - Gently blot excess water and dry the sensor.
 - Recalibrate if instructed to do so by the manufacturer's guidelines.

Most manufacturers recommend recalibration every 4 to 8 hours.

- Remember that response time for gas measurements is slow and values will not always immediately reflect physiologic changes.
 - Average 90% response time for O_2 is 15 to 20 seconds.
 - Average 90% response time for CO_2 is 60 to 90 seconds.

Problem	Technical Solution	Clinical
$P_{tc}O_2 < PaO_2$	1. Recalibrate	1. Presence of shock
1. Improper calibration	2. Allow longer warm-up period	2. Use with high-dose tolazoline", Isuprel, dopamine
2. Insufficient warm-up period after electrode	3. Increase heating temperature	3. Obstructive heart disease with hypoperfusion
application		4. Edema
3. Insufficient heating temperature		5. Severe hypothermia
$P_{tc}O_2 > PaO_2$		
1. Improper calibration	1. Recalibrate	1. Right-to-left ductal shunt with
2. Reading taken immediately after electrode application	y 2. Allow longer warm-up period	preductal electrode and postductal arterial sample
3. Air bubble beneath membrane or leak to atmosphere	 Reapply electrode Attempt calibration at lower temperature 	2. General anesthesia ^{b}
4. Excessive heating temperature	-	

TABLE 9.1 Poor Correlation of PtcO2 and PaO2 Technical Solution Clinical

tc, transcutaneous.

^{*a}</sup><i>In the presence of tolazoline*, P_{tc} O₂ measurements appear to be accurate at lower doses (1–2 mg/kg/h), becoming increasingly inaccurate as dosage increases (43). ^{*b*}Effect depends on concentration and type of membrane.</sup>

I. Complications

- Skin blisters or burns (44, 56, 57)
- Management based on erroneous readings if the unit was not calibrated properly or site precautions were not adhered to (Table 9.1)

Continuous Umbilical Artery PO₂ Monitoring (58, 59)

The following method for monitoring PO_2 and the subsequent method for monitoring blood gases are included for completeness;, however, the editors are not aware of any current commercial source of the required equipment in the United States.

A. Purpose

• Continuous arterial PO₂ monitoring from the umbilical artery

Continuous P_aO_2 monitoring through the umbilical artery offers a means for determining precise data on a continuous basis.

• Trending of P_aO_2 over time

B. Background

- Dual-purpose biluminal catheter
 - A miniature polarographic bipolar oxygen electrode is incorporated into the tip of a bilumen umbilical catheter.
 - \circ The small lumen contains the wires for the electrode.

- The larger lumen can be used for blood sampling, infusion, blood pressure monitoring, and sampling for instrument calibration.
- The electrode is covered by a gas-permeable membrane, under which is a layer of dried electrolyte. The probe is packed dry, and then is activated before use. Water vapor from the activating (hydrating) solution diffuses through the membrane to form a thin layer of liquid electrolyte on the surface of the electrode.
- \circ While it is in the artery, the electrode will produce an electrical current proportional to the PO₂ in the blood.
- $\circ~$ The device is calibrated to the PO_2 value obtained from a blood sample drawn from the catheter.

C. Contraindications

- Previous history of or evidence of compromise to the vascular supply of the lower extremities or the buttock area
- History of previous complications related to an umbilical arterial line
- Peritonitis
- Necrotizing enterocolitis
- Omphalitis
- Omphalocele

D. Equipment

Previously commercially available monitoring systems have been withdrawn from the market recently because of high production costs.

E. Precautions

See also Chapter 28.

- Because this specialized catheter is stiffer and has a wider outer diameter than other umbilical artery catheters, there is the theoretical possibility of a higher rate of failure to insert and vascular spasm and interference with peripheral blood flow.
- Failure to insert this catheter does not imply that insertion of other arterial catheters will be unsuccessful.
- The electrode may fail to activate or may lose activation.
- The catheter should be removed slowly to ensure that physiologic vasospasm occurs with removal.

F. Technique

- Use sterile procedure.
- Prepare the catheter according to the manufacturer's instructions.
- 4 Fr catheters are recommended for infants weighing less than 1,500 g.
- The technique for placement/insertion is the same as that used for the placement of conventional umbilical artery catheters. Either a high or low umbilical artery placement can be used (see Chapter 28).
- Verify catheter position by radiography.
- Draw blood sample for calibration.
- Calibrate the monitor according to the manufacturer's instructions.

G. Complications

Same as for umbilical artery catheterization. See Chapter 28.

Continuous Umbilical Artery PO₂, PCO₂, pH, and Temperature Blood Gas Monitoring (60, 61, 62, 63, 64 and 65)

A. Purpose

• Continuous arterial blood gas monitoring from the umbilical artery

Continuous blood gas monitoring through the umbilical artery offers a means for determining precise data on a continuous basis.

• Trending of blood gas data over time

B. Background

- A very thin, multiparameter, single-use disposable fiber-optic sensor
 - Measures pH, PCO₂, PO₂, and temperature directly
 - Introduced into the bloodstream via the umbilical artery catheter
 - Port allows blood sampling, blood pressure monitoring, and drug infusion
- Calculated parameters include bicarbonate, base excess, and oxygen saturation.
- Delivers continuous ventilation, oxygenation, and acid balance information, while also conserving blood volume by reducing blood sampling

C. Contraindications

- Previous history or evidence of compromise to the vascular supply of the lower extremity or the buttock area
- History of previous complications related to an umbilical arterial line
- Peritonitis
- Necrotizing enterocolitis
- Omphalitis
- Omphalocele

D. Equipment

Previously commercially available monitoring systems have been withdrawn from the market recently because of high production costs.

E. Precautions

- The fiber-optic sensor may fail as a result of excessive kinking during sensor insertion into the umbilical artery catheter.
- The sensor should be removed slowly to ensure that there is no microthrombus release if heparinization of the catheter was suboptimal.
- See also Chapter 28.

F. Technique

- Use sterile procedure.
- Insert umbilical artery catheter (see Chapter 28).
- Verify catheter position by radiography.
- Calibrate sensor following the manufacturer's instructions.
- Introduce the sensor into the umbilical artery catheter following the manufacturer's instructions.

G. Complications

Same as for umbilical artery catheterization; see Chapter 28.

References

1. Hay WW. Physiology of oxygenation and its relation to pulse oximetry in neonates. J Perinatol. 1987;7:309.

- 2. Dziedzic K, Vidyasagar D. Pulse oximetry in neonatal intensive care. Clin Perinatol. 1989;16:177.
- 3. Barrington KJ, Finer NN, Ryan CA. Evaluation of pulse oximetry as a continuous monitoring technique in the neonatal intensive care unit. Crit Care Med. 1988;16:1147.
- 4. Solimano AJ, Smyth JA, Mann TK, et al. Pulse oximetry advantages in infants with bronchopulmonary dysplasia. Pediatrics. 1986;78:844.
- 5. Bowes WA, Corke BC, Hulka J. Pulse oximetry: a review of the theory, accuracy and clinical applications. Obstet Gynecol. 1989;74:541.
- 6. Sendak MJ, Harris AP, Donham RT. Use of pulse oximetry to assess arterial oxygen saturation during newborn resuscitation. Crit Care Med. 1986;14:739.
- 7. House JT, Schultetus RR, Graverstein N. Continuous neonatal evaluation in the delivery room by pulse oximetry. J Clin Monit. 1987;96:96.
- 8. Porter KB. Evaluation of arterial oxygen saturation of the newborn in the labor and delivery suite. J Perinatol. 1987;7:337.
- 9. Deckardt R, Schneider K-T, Graeff H. Monitoring arterial oxygen saturation in the neonate. J Perinat Med. 1987;15:357.
- 10. Maxwell LG, Harris AP, Sendak MJ, et al. Monitoring the resuscitation of preterm infants in the delivery room using pulse oximetry. Clin Pediatr. 1987;26:18.
- 11. Ramanathan R, Durand M, Larrazabal C. Pulse oximetry in very low birth weight infants with acute and chronic lung disease. Pediatrics. 1987;79:612.
- 12. Deckardt R, Stewart DJ. Noninvasive arterial hemoglobin oxygen saturation versus transcutaneous oxygen tension monitoring in the preterm infant. Crit Care Med. 1984;12:935.
- 13. Miyasaka K, Katayama M, Kusakawa I, et al. Use of pulse oximetry in neonatal anesthesia. J Perinatol. 1987;7:343.
- 14. Fanconi S. Reliability of pulse oximetry in hypoxic infants. J Pediatr. 1988;112:424.
- 15. Lewallen PK, Mamme MC, Coleman JM, et al. Neonatal transcutaneous oxygen saturation monitoring. J Perinatol. 1987;7:8.
- 16. Praud JP, Carofilis A, Bridey F, et al. Accuracy of two wavelength pulse oximetry in neonates and infants. Pediatr Pulmonol. 1989;6:180.
- 17. Southall DP, Bignall S, Stebbens VA, et al. Pulse oximeter and transcutaneous arterial oxygen measurements in neonatal and pediatric intensive care. Arch Dis Child. 1987;62:882.
- 18. Hay WW. The uses, benefits and limitations of pulse oximetry in neonatal medicine: consensus on key issues. J Perinatol. 1987;7:347.
- 19. Emery JR. Skin pigmentation as an influence on the accuracy of pulse oximetry. J Perinatol. 1987;7:329.
- 20. Durand M, Ramanathan R. Pulse oximetry for continuous oxygen monitoring in sick newborn infants. J Pediatr. 1986;109:1052.
- 21. Jennis MS, Peabody JL. Pulse oximetry: an alternative method for the assessment of oxygenation in newborn infants. Pediatrics. 1987;79:524.
- 22. Wasunna A, Whitelaw GL. Pulse oximetry in preterm infants. Arch Dis Child. 1987;62:957.
- 23. Reynolds GJ. Guidelines for the use of pulse oximetry in the noninvasive estimation of oxygen saturation in oxygen-dependent newborn infants. Aust Paediatr J. 1988;24:346.
- 24. Anderson JV. The accuracy of pulse oximetry in neonates: effects of fetal hemoglobin and bilirubin. J Perinatol. 1987;7:323.
- 25. Bucher H-U, Fanconi S, Baeckert P, et al. Hyperoxemia in newborn infants: detection by pulse oximetry. Pediatrics. 1989;84:226.
- 26. Hay WW, Brockway J, Eyzaguirre M. Neonatal pulse oximetry: accuracy and reliability. Pediatrics. 1989;83:717.
- 27. Kopotic RJ, Mannino FL, Colley CD, et al. Display variability, false alarms, probe cautions, and recorder use in neonatal pulse oximetry. J Perinatol. 1987;7:340.

28. Sobel DB. Burning of a neonate due to a pulse oximeter: arterial saturation monitoring. Pediatrics. 1992;89:154.

29. Ghat R, Diaz-Blanco J, Chaudhry U, et al. Recent instrumentation. Pediatr Clin North Am. 1986;33:503. 30. Huch R, Huch A, Albani M, et al. Transcutaneous PO_2 monitoring in routine management of infants and children with cardiorespiratory problems. Pediatrics. 1976;57:681.

31. Okken A, Rubin IL, Martin RJ. Intermittent bag ventilation of preterm infants on continuous positive airway pressure: the effect on transcutaneous PCO₂. J Pediatr. 1978;93:279.

32. Hansen TH, Tooley WH. Skin surface carbon dioxide tension in sick infants. Pediatrics. 1979;64:942.33. Laptook A, Oh W. Transcutaneous carbon dioxide monitoring in the newborn period. Crit Care Med. 1981;9:759.

34. Monaco F, McQuitty J. Transcutaneous measurement of carbon dioxide partial pressure in sick neonates. Crit Care Med. 1981;9:756.

35. Clarke TA, Zmora E, Chen J-H, et al. Transcutaneous oxygen monitoring during neonatal transport. Pediatrics. 1980;65:884.

36. O'Connor TA, Grueber R. Transcutaneous measurement of carbon dioxide tension during long-distance transport of neonates receiving mechanical ventilation. J Perinatol. 1998; 18:189

37. Beachy P, Whitfield J. The effect of transcutaneous PO₂ monitoring on the frequency of arterial bloodgas analysis in the newborn with respiratory distress. Crit Care Med. 1981; 9:584.

38. Slavin RE, Cohen A, Epstein MF, et al. Two-site noninvasive oxygen monitoring in a case of persistent fetal circulation. Respir Care. 1980;25:358.

39. Tateishi K, Yamanouchi I. Noninvasive transcutaneous oxygen pressure diagnosis of reversed ductal shunts in cyanotic heart disease. Pediatrics. 1980;66:22.

40. Versmold HT, Linderkamp O, Holzman M, et al. Limits of TcPO₂ monitoring in sick neonates: relation to blood pressure, blood volume, peripheral blood flow, and acid base status. Acta Anaesthesiol Scand. 1978;68:88.

41. Beran A, Tolle C, Huxtable R. Cutaneous blood flow and its relationship to transcutaneous O_2/CO_2 measurements. Crit Care Med. 1981;9:736.

42. Rowe MI, Weinberg G. Transcutaneous oxygen monitoring in shock and resuscitation. J Pediatr Surg. 1979;14:773.

43. Peabody JL. Transcutaneous oxygen measurement to evaluate drug effects. Clin Perinatol. 1979;6:109. 44. Avery GB, Bancalari EH, Engler A, et al. Task Force on Transcutaneous Oxygen Monitors: report of consensus meeting December 5 to 6, 1986. Pediatrics. 1989;83:122.

45. Peabody JL. Historical perspective of noninvasive monitoring. J Perinatol. 1987;7:306.

46. Fanconi S, Sigrist H. Transcutaneous carbon dioxide and oxygen tension in newborn infants: reliability of a combined monitor of oxygen tension and carbon dioxide tension. J Clin Monit. 1988;4:103.

47. Yip WCL, Ho TF, Tay JSH, et al. The application of transcutaneous oxygen monitoring in paediatric intensive care: a critical appraisal of reliability and safety. J Singapore Paediatr Soc. 1983;25:33.

48. Fanconi S, Doherty P, Edmonds JF, et al. Pulse oximetry in pediatric intensive care: comparison with measured saturations and transcutaneous oxygen tension. J Pediatr. 1985;107:362.

49. Kraus AN, Waldman S, Frayer W, et al. Noninvasive estimation of arterial oxygenation in newborn infants. J Pediatr. 1978;93:275.

50. Rooth G, Huch A, Huch R. Transcutaneous oxygen monitors are reliable indicators of arterial oxygen tension (if used correctly). Pediatrics. 1987;79:283.

51. Rome ES, Stork EK, Carlo WA, et al. Limitations of transcutaneous PO_2 and P_aO_2 measurements in infants with BPD. Pediatrics. 1984;74:217.

52. Solimano AJ, Smyth JA, Mann TK, et al. Pulse oximetry advantages in infants with bronchopulmonary dysplasia. Pediatrics. 1986;78:844.

53. Mok JYQ, McLaughlin FJ, Pintar M, et al. Transcutaneous monitoring of oxygenation: what is normal? J Pediatr. 1986;108:365.

54. Palmisano BW, Severinghaus JW. Transcutaneous PCO₂ and PO₂: a multicenter study of accuracy. J Clin Monit. 1990;6:189.

55. Pearlman SA, Maisels MJ. Preductal and postductal transcutaneous oxygen tension measurements in premature newborns with hyaline membrane disease. Pediatrics. 1989;83:98.

56. Golden SM. Skin cratersâ€"a complication of transcutaneous oxygen monitoring. Pediatrics. 1981;67:514.

57. Wimberley PD, Frederiksen PS, Witt-Hansen PS, et al. Evaluation of a transcutaneous oxygen and carbon dioxide monitor in a neonatal intensive care department. Acta Paediatr Scand. 1985;74:352. 58. Fink SE. Continuous P_aO_2 monitoring through the umbilical artery. Neonat Intensive Care. 1990;3:16. 59. Phillips BL, Quitty J, Durand DJ. Blood gases: technical aspects and interpretation. In: Goldsmith JP, Karotkin EH, Barber S, eds. Assisted Ventilation of the Neonate. 2nd ed. Philadelphia: Saunders; 1988:213. 60. Weiss I, Fink S, Harrison R, et al. Clinical use of continuous arterial blood-gas monitoring in the pediatric intensive care unit. Pediatrics. 1999;103:440.

61. Coule LW, Truemper EJ, Steinhart CM, et al. Accuracy and utility of a continuous intra-arterial blood-gas monitoring system in pediatric patients. Crit Care Med. 2001;29:420.

62. Meyers PA, Worwa C, Trusty R, Mammel MC. Clinical validation of a continuous intravascular neonatal blood gas sensor introduced through an umbilical artery catheter. Respir Care. 2002;47(6):682.
63. Rais-Bahrami K, Rivera O, Mikesell GT, Short BL. Continuous blood gas monitoring using an indwelling optode method: comparison to intermittent arterial blood gas sampling in ECMO patients. J Perinatol. 2002;22(6):472.

64. Rais-Bahrami K, Rivera O, Mikesell GT, Short BL. Continuous blood gas monitoring using an indwelling optode method: clinical evaluation of the Neotrend® sensor using a Luer stub adaptor to access the umbilical artery catheter. J Perinatol. 2002; 22(5):367.

65. Ganter M, Zollinger A. Continuous intravascular blood gas monitoring: development, current techniques, and clinical use of a commercial device. Br J Anaesth. 2003;91(3):397. Review.

10-Capnography

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A. Definitions

Capnography refers to the continuous analysis and recording of carbon dioxide concentrations in respiratory gases ($P_{et}CO_2$), whereas capnometry refers to measurement or analysis alone of carbon dioxide concentrations without a continuous written record or waveform. A device that measures and displays the breath-to-breath numeric values of CO_2 is referred to as capnometer, whereas a device that also displays the waveform of CO_2 during the respiratory cycle is called a capnograph. B. Purpose

- Noninvasive continuous analysis and recording of CO₂ during tidal breathing (1,2 and 3)
- End-tidal CO₂-level monitoring (P_{et}CO₂)
- Respiratory rate monitoring

C. Background

- Most capnography employs infrared technology, which is based on the infrared absorption of CO₂.
 - Capnographic devices incorporate one of two types of analyzers: mainstream and sidestream.
 - With a mainstream analyzer, the sensor is attached directly to an optical adapter that is inline with the endotracheal tube (Fig. 10.1).
 - With a sidestream analyzer, a low-dead-space adapter is placed inline with the endotracheal tube and gas is aspirated continuously to the analyzer for measurement (Fig. 10.2).
 - For both mainstream and sidestream capnography, the only adapters that should be used for newborns are ones specifically designed for neonatal application.

D. Indications

- Evaluation of the exhaled CO₂, specifically end-tidal CO₂, which is the maximum partial pressure of CO₂ exhaled during a tidal breath just prior to the beginning of inspiration (designated P_{et}CO₂) (4,5) (Fig. 10.2)
- Monitoring the severity of pulmonary disease and evaluating response to therapy, particularly therapy intended to change the ratio of dead space to tidal volume (6) or to improve the matching of ventilation to perfusion (V/Q) (7)
- Determining that tracheal rather than esophageal intubation has taken place (low or absent cardiac output may negate its use for this indication)
- Continued monitoring of the integrity of the ventilatory circuit (8)
- More accurate and continuous reflection of CO_2 elimination (9,10)
- Graphic evaluation (11)
- Use of capnography in combination with pulse oximetry can allow for additional monitoring to detect airway obstruction or subclinical degrees of respiratory depression in the sedated patient (12).
- Evaluation of surfactant efficacy can be augmented by use of dead-space and capnography measurements (13).
- Sidestream technology can also be used with nasal prongs in spontaneously breathing patients.

E. Contraindications

There are no absolute contraindications to capnography in the mechanically ventilated infant.

F. Limitations (10,14 15,16 and 17)

- The composition of the respiratory gas mixture may affect the capnogram; the infrared spectrum of CO₂ has some similarities to the spectra for both oxygen and nitrous oxide (most available capnographs have a correction factor already incorporated into the calibration).
- Rapid changes in respiratory rate and tidal volume may lead to measurement error, depending on the frequency response of the capnograph; different capnographs may have different frequency responses.
- Contamination of either the monitor or the sampling system by secretions, blood, or condensation may lead to inaccurate results.
- Large dead space affects P_{et}CO₂ measurements. The difference between P_{et}CO₂ and PaCO₂ increases as dead-space volume increases and may vary within the same patient over time.
- The end-tidal CO₂ adapter can add to the dead space and resistance of the respiratory circuit, particularly in small infants.
- P_{et}CO₂ measurements may not provide an accurate correlation with PaCO₂ in small preterm infants with nonhomogenous lung disease and therefore cannot be substituted for PaCO₂ analyses in preterm infants during this critical period (18,19).



FIG. 10.1. Mainstream capnographic monitor unit and clip-on sensor assembly. (Courtesy of Respironics, Inc., Murrysville, PA, USA.)



FIG. 10.2. Sidestream capnographic monitor unit with sample line. Note end-tidal CO_2 value and wave. (Courtesy of Oridion Capnography, Needham, MA, USA.)

G. Equipment

- For mainstream capnography, an airway adapter is needed, along with a reusable sensor attachment.
- For sidestream capnography, an airway adapter with sampling tube is used (Fig. 10.3).
- Capnogram



FIG. 10.3. Infant sidestream lowâ€"dead-space adapter with sample tubing. (Courtesy of Oridion Capnography, Needham, MA, USA.) H. Precautions

- In the mainstream adapter, prevent condensation in the airway adapter.
- In the sidestream adapter, prevent fluid (water) buildup in the sample tube.
- For both mainstream and sidestream, when adding bulk to the endotracheal tube, more attention should be given to properly securing the position of the tube.
- Tidal volume measurements may be affected if the end-tidal CO_2 adapter is placed between the endotracheal tube and the ventilator flow sensor.



FIG. 10.4. Infant sidestream lowâ€"dead-space adapter (see arrow) inline with endotracheal tube.

I. Technique

- Familiarize yourself with the system before proceeding.
- Attach the adapter inline with the endotracheal tube and the ventilator T piece (both sidestream and mainstream) (Fig. 10.4).
- For mainstream capnography, connect the sensor to the airway adapter.
- For sidestream capnography, connect the sampling tube to the analyzer.

J. Complications

Capnography is a safe, noninvasive test, with some limitations, depending on the type of capnography that is employed (2).

- With mainstream analyzers, the use of too large an airway tube adapter together with the weight of the probe may introduce an excessive amount of bulk and weight to the endotracheal tube.
- With sidestream capnography, a lowâ€"dead-space adapter allows for less bulk and weight; however, care must be taken not to pull excessively on the sample line that is going to the measurement instrument.

References

1. Campbell RS, Branson RD, Burke W, et al. Capnography/capnometry during mechanical ventilation. Respir Care. 1995; 40:1321.

2. Hess DR, Branson RD. Noninvasive respiratory equipment. In: Branson RD, Hess DR, Chatburn RL, eds. Respiratory Care Equipment. Philadelphia: Lippincott; 1994:184.

3. Block FE, McDonald JS. Sidestream versus mainstream carbon dioxide analyzers. J Clin Monit. 1992;8:139.

4. Carlon GC, Ray C, Miodownik S, et al. Capnography in mechanically ventilated patients. Crit Care Med. 1988;16:550.

5. Gravenstein N, Good ML. Noninvasive assessment of cardiopulmonary function. In: Civetta JM, Taylor RW, Kirby RR, eds. Critical Care. Philadelphia: Lippincott; 1988:291.

6. Yamanaka MK, Sue DY. Comparison of arterial-end-tidal PCO₂ difference and dead space/tidal volume ratio in respiratory failure. Chest. 1987;92:254.

7. Burrows FA. Physiologic dead space, venous admixture, and the arterial to end-tidal carbon dioxide difference in infants and children undergoing cardiac surgery. Anesthesiology. 1989;70:219.

8. Eichhorn JH, Cooper JB, Cullen DJ, et al. Standards for patient monitoring during anesthesia at Harvard Medical School. JAMA. 1986;256:1017.

9. Morley TF, Giamo J, Maroszan E, et al. Use of capnography for assessment of the adequacy of alveolar ventilation during weaning from mechanical ventilation. Am Rev Respir Dis. 1993;148:339.

10. Rozycki HJ, Sysyn GD, Marshall MK, et al. Mainstream end-tidal carbon dioxide monitoring in the neonatal intensive care unit. Pediatrics. 1998;101:648.

11. Bhavani-Shankar K, Moseley H, Kumar AY, et al. Capnography and anesthesia. Can J Anesth. 1992;39:617.

12. Tobias J. End-tidal carbon dioxide monitoring during sedation with a combination of midazolam and ketamine for children undergoing painful, invasive procedures. Pediatr Emerg Care. 1999;15:173.

13. Wenzel U, Rudiger M, Wagner M, et al. Utility of deadspace and capnometry measurements in determination of surfactant efficacy in surfactant-depleted lungs. Crit Care Med. 1999;27:946.

14. Graybeal JM, Russell GB. Capnometry in the surgical ICU: an analysis of the arterial-to-end-tidal carbon dioxide difference. Respir Care. 1993;38:923.

15. Hess D. Capnometry and capnography: technical aspects, physiologic aspects, and clinical applications. Respir Care. 1990;35:557.

16. Isert P. Control of carbon dioxide levels during neuroanesthesia: current practice and an appraisal of our reliance upon capnography. Anaesth Intensive Care. 1994;22:435.
17. Saili N, Dutta AK. End tidal carbon dioxide monitoringâ€"its reliability in neonates. Indian J Pediatr. 1997;64:389.

18. Aliwalas LL, Noble L, Nesbitt K, et al. Agreement of carbon dioxide levels measured by arterial, transcutaneous and end tidal methods in preterm infants < or = 28 weeks gestation. J Perinatol. 2005;25(1):26.

19. Tingay DG, Stewart MJ, Morley CJ. Monitoring of end tidal carbon dioxide and transcutaneous carbon dioxide during neonatal transport. Arch Dis Child Fetal Neonatal Ed. 2005;90(6):F523.

11-Transcutaneous Bilirubin Testing

Aimee M. Barton

A. Background

- Jaundice occurs in most newborn infants. Bilirubin at high levels is potentially toxic to the nervous system, causing bilirubin encephalopathy and kernicterus (1).
- A systematic assessment of all newborn infants for the risk of severe hyperbilirubinemia should be undertaken prior to discharge, and appropriate follow-up should be provided. This assessment may be performed by measuring total serum bilirubin (TSB) or transcutaneous bilirubin (TCB) (2).
- Visual assessment of jaundice, although clinically important, may not be accurate (3, 4).
- Transcutaneous bilirubinometers measure the yellowness of reflected light from the skin and subcutaneous tissues to provide an objective noninvasive measurement of the degree of neonatal jaundice and thereby predict the approximate TSB.
- Transcutaneous bilirubinometers are used predominantly for screening for significant hyperbilirubinemia in term and near-term newborn infants.
- Two transcutaneous bilirubinometers are currently in use in the United States. Although these instruments use different technologies and algorithms, their underlying principles of operation are similar.
 - Konica Minolta/Air-Shields JM-103 Jaundice Meter (Dräger Medical, Telford, PA, USA) (4, 5) (Fig. 11.1)
 - BiliChek Noninvasive Bilirubin Analyzer (Children's Medical Ventures/Respironics, Norwell, MA, USA) (4, 6) (Fig. 11.2)
- Both bilirubinometers have been found to provide TCB measurements that correlate well with TSB values at levels <15 mg/dL, in term and near-term newborn infants, but wider variations have been noted at higher bilirubin levels (6, 7).

B. Indications

- TCB may be obtained:
 - As part of routine predischarge assessment between 1 and 4 days of life in term and near-term newborn infants, to assess the risk of development of severe hyperbilirubinemia, by using the hour-specific bilirubin nomogram (2, 8). (Fig. 11.3)
 - For repeated noninvasive measurement of progression of jaundice in term or near-term newborn infants.
 - When clinical jaundice is noted in the first 24 hours of life
 - When jaundice appears excessive for the infant's age
- TSB (in addition to other studies to determine underlying pathology) should be obtained when (2):
 - Infant is receiving phototherapy or the TSB is rising rapidly
 - TSB concentration approaching exchange transfusion levels or not responding to phototherapy

- Infant has an elevated direct bilirubin level
- Jaundice is present at or beyond age 3 weeks
- In sick or premature (<35 weeks' gestation) infants

C. Limitations

- TCB measurement is a screening tool and should not be used for treatment decisions, but rather to select those infants who should undergo TSB measurement (1).
- The two large studies evaluating the BiliChek device and the JM-103 device included few patients with TSB values >15 mg/dL. The accuracy of TCB measurement in this range has not been evaluated adequately (6, 7).
- TCB measurements become less accurate if the infant is being treated with phototherapy or has received an exchange transfusion and should not be used within 24 hours of either of these therapies (2, 4, 9, 10).
 - Phototherapy alters the chemical structure of bilirubin in the subcutaneous tissues, making it more water-soluble. Measurement of TCB in infants undergoing phototherapy is not reliable because the large decrease in subcutaneous bilirubin may not yet be reflected in the serum (9, 10).
 - One study found that correlation coefficients decreased from 0.90 before phototherapy to 0.85 upon initiation of phototherapy. This trend continued with prolonged phototherapy, with correlation coefficients as low as 0.33 in infants undergoing phototherapy for longer than 48 hours (9).
- TCB measurements are less accurate in preterm infants.
 - When compared with TSB measurement, the BiliChek System has been shown to have a correlation coefficient of 0.86 in preterm infants who are not receiving phototherapy (10).



FIG. 11.1. Use of the Konica Minolta/Air-Shields JM-103 Jaundice Meter on the sternum. (Photo provided by Dräger Medical.)



FIG. 11.2. Use of the BiliChek Noninvasive Bilirubin Analyzer on the forehead. (Photo provided by Children's Medical Ventures/Respironics.)



FIG. 11.3. Nomogram for designation of risk in 2,840 well newborns at 36 or more weeks' gestational age with birth weight of 2,000 g or more or 35 or more weeks' gestational age and birth weight of 2,500 g or more based on hour-specific serum bilirubin values (2, 8). (Reproduced with permission from Pediatrics, Vol. 114, Page 301, Copyright \hat{A} [©] 2004 by the AAP.)

D. Equipment

- Two TCB monitors are currently in use in the United States
 - Konica Minolta/Air-Shields JM-103 Jaundice Meter (Dräger Medical, Telford, PA, USA) (Figs. 11.1 and 11.4)

- Determines the yellowness of subcutaneous tissue by measuring the difference between optical densities for light in the blue and green wavelengths
- Measurement probe has two optical paths
- By calculating the difference between the optical densities, the parts common to the epidermis and dermis are deducted. As a result, the difference can be obtained for subcutaneous tissue only.
- Theoretically allows for measurement of degree of yellowness of skin and subcutaneous tissue with minimal influence of melanin pigment and skin maturity
- Linear correlation of this measurement with TSB allows for conversion to TSB by the meter, which is indicated digitally.
- BiliChek Noninvasive Bilirubin Analyzer (Children's Medical Ventures/Respironics, Norwell, MA, USA) (Figs. 11.2 and 11.5)
 - Noninvasive device consisting of light source, microspectrophotometer, fiber-optic probe, and microprocessor control circuit
 - Uses entire spectrum of visible light reflected by the skin
 - White light is transmitted into the skin and the reflected light is collected for analysis.
 - Algorithms take into account the effect of hemoglobin, melanin, and dermal thickness.
 - Absorption of light due to bilirubin in the capillary bed and subcutaneous tissue is isolated by spectral subtraction.
- E. Special Circumstances/Considerations
 - Hospital protocols should include the conditions under which TCB and TSB levels are to be obtained (1).
 - Only TSB measurements should be performed in infants with severe enough jaundice to warrant exchange transfusion (6).
 - TCB is less accurate in infants undergoing phototherapy and therefore serum levels are preferred for monitoring bilirubin values in such infants (2, 4, 8, 9).
 - Race/skin color: TCB readings obtained by the BiliChek have been found to correlate with TSB values in white, black, Asian, Hispanic, indigenous African, and Indian infants (6, 9, 11). In black infants, TCB readings obtained by the JM-103 correlate less closely with TSB values, with the TCB generally being greater than the TSB (7).



JM-103

FIG. 11.4. Measurement principle of Konica Minolta/Air-Shields JM-103 Jaundice Meter. (Reproduced with permission from

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F. Techniques

- Calibrate the TCB device according to manufacturer specifications.
- Measure TCB by pressing the trigger button and gently pressing the tip to the infant's forehead or sternum until the device indicates that reading is complete.
 - Some studies have shown that TCB measurements from the sternum correlated slightly better 0 with TSB levels than TCB measurements from the forehead, possibly as a result of the exposure of the forehead to ambient light. Other studies indicate both sites to be equivalent (7, 12).
 - Measurements must be taken in a consistent manner with regard to placement of the probe 0 and amount of pressure applied to the device. Interoperative and intraoperative variability may be minimized with proper training (7).

- Measurement of the TCB using the BiliChek system takes approximately 20 to 80 seconds. This time is required for the monitor to make five measurements that are averaged to provide one TCB value. The JM-103 takes approximately 10 seconds to obtain its dual measurements and calculate the TCB value.
- Repeated use of the disposable probes is not recommended.



FIG. 11.5. Measurement principle of the BiliChek Noninvasive Bilirubin Analyzer. (Reproduced with permission of Children's Medical Ventures/Respironics.) G. Complications

• No complications have been reported from the use of TCB monitors, apart from the risk of inappropriate use and the possibility of underestimation of the level of jaundice.

References

1. American Academy of Pediatrics Subcommittee on Hyperbilirubinemia. Neonatal jaundice and kernicterus. Pediatrics. 2001;108(3):31–39.

 American Academy of Pediatrics Subcommittee on Hyperbilirubinemia. Management of hyperbilirubinemia in the newborn infant 35 or more weeks of gestation. Pediatrics. 2004;114:297–316.
 Szabo P, Wolf M, Bucher HU, et al. Detection of hyperbilirubinaemia in jaundiced full-term neonates by eye or bilirubinometer? Eur J Pediatr. 2004;163(12):722–727.

4. Maisels MJ. Transcutaneous bilirubinometry. NeoReviews. 2006;7(5):e217–e225.

5. Yasuda S, Itoh S, Isobe K, et al. New transcutaneous jaundice device with two optical paths. J Perinat Med. 2003;31:81–88.

6. Bhutani VK, Gourley GR, Adler S, et al. Noninvasive measurement of total serum bilirubin in a multiracial predischarge newborn population to assess the risk of severe hyperbilirubinemia. Pediatrics. 2000;106:e17.

7. Maisels MJ, Ostrea EM, Touch S, et al. Evaluation of a new transcutaneous bilirubinometer. Pediatrics. 2004;113:1628–1635.

8. Bhutani VK, Johnson L, Sivieri EM. Predictive ability of a predischarge hour-specific serum bilirubin for subsequent significant hyperbilirubinemia in healthy term and near-term newborns. Pediatrics. 1999;103(1):6–14.

9. Mahajan G, Kaushal RK, Sankhyan N, et al. Trancutaneous bilirubinometer in assessment of neonatal jaundice in Northern India. Indian Pediatr. 2005;42:41–45.

10. Nanjundaswamy S, Petrova A, Mehta R, Hegyi T. Transcutaneous bilirubinometry in preterm infants receiving phototherapy. Am J Perinatol. 2005;22(3):127–131.

11. Slusher TM, Angyo IA, Bode-Thomas F, et al. Transcutaneous bilirubin measurements and serum total bilirubin levels in Indigenous African infants. Pediatrics. 2004;113:1636–1641.

12. Ebbesen F, Rasmussen LM, Wimberley PD. A new transcutaneous bilirubinometer, BiliChek®, used in the neonatal intensive care unit and the maternity ward. Acta Paediatr. 2002; 91:203–211.

12-Vessel Transillumination

Dawn M. Walton Billie Lou Short

A. Indications (1,2,3,4, and 5)

- To locate artery or vein for:
 - Puncture for sampling
 - Vessel cannulation
 - \circ See also Chapter 36, for use in diagnosing thoracic air leaks (1,2,3,4 and 5).

B. Contraindications None C. Precautions

C. Flecautions

- Use fiber-optic light source with appropriate filters to cool light and prevent burns (6,7).
- Place sterile surgical glove over tip of probe to preserve sterile field.

D. Equipment

- Transillumination source
 - High-intensity fiber-optic light (black soft tube around probe can make light seal)
 - Otoscope light may be used in some instances (8).
- Alcohol
- Sterile glove

E. Technique

- Clean end of light source with an alcohol swab. Cover with sterile glove.
- Dim light in room. Some residual light is necessary to visualize operating field.
- Set light source at low intensity and increase as needed for visualization.
- Position probe to transilluminate vessel.
 - Directly opposite to puncture site, through extremity
 - Adjacent to vessel but out of way of procedure
- Identify vessel as dark, linear structure (Fig. 12.1).

- Edges may be indistinct.
- Arteries will be pulsatile.
- Compensate for distortion if light is not directly opposite puncture site.
- Do not maintain contact between light source and extremity for long periods of time.

F. Complications

- Burns from light probe (Fig. 12.2) (7)
- Cross-contamination from breach of sterile technique



FIG. 12.1. Transillumination. A: Arteries and veins on volar aspect of left wrist. B: Venous arch on dorsum of hand. C: Vessels in right antecubital fossa. D: Left posterior tibial artery. P.83



FIG. 12.2. Burn from transilluminator.

References

1. Cole FS, Todres ID, Shannon DC. Technique for percutaneous cannulation of the radial artery in the newborn infant. J Pediatr. 1978;92:105.

2. Curran JS, Ruge W. A restraint and transillumination device for neonatal arterial/venipuncture: efficacy and thermal safety. Pediatrics. 1980;66:128.

3. Mattson D, O'Connor M. Transilluminator assistance in neonatal venipuncture. Neonatal Network. 1986;5:42.

4. Schwartz N, Eisenkraft JB. Transillumination aids the percutaneous cannulation of peripheral vessels. J Cardiothorac Anesth. 1989;3:675.

5. Dinner M. Transillumination to facilitate venipuncture in children. Anesth Analg. 1992;74:467.

6. Sajben FP, Gibbs NF, Friedlander SF. Transillumination blisters in a neonate. J Am Acad Dermatol. 1999;41:264.

7. Keroack MA, Kotilainen HR, Griffin BE. A cluster of atypical skin lesions in well-baby nurseries and a neonatal intensive care unit. J Perinatol. 1996;16:370.

8. Goren A, Laufer J, Yativ N, et al. Transillumination of the palm for venipuncture in infants. Pediatr Emerg Care. 2001; 17(2):130.

13-Venipuncture

Dawn M. Walton Billie Lou Short

A. Indications

- Blood sampling
 - Routine, particularly if a large volume of blood is needed
 - Blood culture
 - Central hematocrit
 - Preferred (over capillary sample) for certain studies (1,2 and 3)
 - Ammonia (arterial optimal)
 - Drug levels
 - Cross-matching blood
 - Hemoglobin/hematocrit
 - Karyotype
 - Lactate and pyruvate levels (arterial optimal)
- Administration of drugs

B. Contraindications

- Use of deep vein in presence of coagulation defect
- Local infection at puncture site
- Femoral or internal jugular vein (see G)
- External jugular vein in infants with respiratory distress, intracranial hemorrhage, or raised intracranial pressure

C. Precautions

- Observe universal precautions. Wear gloves.
- When sampling from neck veins, place infant in head-down position to avoid cranial air embolus. Do not use neck veins in infants with intracranial bleeding or increased intracranial pressure, except as a last resort.
- Remove tourniquet before removing needle, to minimize hematoma formation.
- Apply local pressure with dry gauze to produce hemostasis (usually 2 to 3 minutes).
- Avoid using alcohol swab to apply local pressure (painful, impairs hemostasis).

D. Special Considerations for Neonates

- Conserve sites to preserve limited venous access by using distal sites first whenever possible.
- Use small needle or scalp vein butterfly. A 23-gauge needle is best. Hemolysis or clotting may occur with a 25 gauge or smaller.
- Choice of veins (Fig. 13.1) in order of preference:
 - Antecubital fossa
 - Dorsum of hands
 - Dorsum of feet
 - Greater saphenous vein at the ankle
 - Vein in center of the volar aspect of the wrist
 - o Scalp
 - Proximal greater saphenous vein

- o Neck
- Recent studies show that adequate pain control can be achieved during venipuncture with EMLA (Astra Pharmaceuticals, L.P., Wayne, PA, USA) cream applied 1 hour prior to procedure, if time allows (4,5).
- Oral sucrose solution (24% to 25%) provides quick and effective pain control for venipuncture (6,7).
- Heel lancing can be more painful and require more punctures than venipuncture in infants (4,8).

E. Equipment

- Gloves
- A 23- to 25-gauge venipuncture needle (a safety-engineered needle should be used) (Fig. 13.2).
- Syringe with volume just larger than sample to be drawn
- Prepared alcohol swabs
- Gauze pads
- Appropriate containers for specimens
- For blood culture:
 - Povidone-iodine solution preparation (three swabs)
 - Sterile gloves
 - Blood culture bottle(s)
 - Transfer needle
- Tourniquet or sphygmomanometer cuff







FIG. 13.2. Safety-engineered needles for venipuncture.

F. Technique (See Procedures DVD for Video) General Venipuncture

- Locate the appropriate vessel. Use transillumination if necessary (see Chapter 12). Warm extremity with heel warmer or warm washcloth if circulation is poor.
- Apply anesthetic cream if time permits, and/or administer sucrose solution if possible.
- Restrain infant appropriately.

•

- Prepare area with antiseptic (see Chapter 4).
 - Occlude vein proximally using either:
 - Tourniquet or cuff inflated to level between systolic and diastolic pressure (Fig. 13.3)
 - Direct pressure over vessel
 - Rubber band (loop two bands together, tied as in Fig. 13.3)





FIG. 13.4. Anterior wall of vein removed. Needle penetrating skin a short distance from site of venipuncture.

- Remove occlusion device and replace to promote optimal vein distension.
- Check syringe function and attach to needle. Alternative method is to use Microlance needle (0.9 x 40 mm) (Becton Dickinson, Franklin Lakes, NJ, USA) without a syringe to collect samples by drip method. Drip method cannot be used for blood culture or coagulation studies (8).
- Penetrate skin first and position for entry of vein (Fig. 13.4).
 - Angle of entry 25 to 45 degrees
 - Bevel up preferred for optimal blood flow (less chance of needle occlusion by vein wall)
 - o Direction of entry with or against direction of blood flow
 - o If possible, insert needle at area where vessel bifurcates to avoid "rolling†of veins.

- Collect sample by gentle suction
 - To prevent occlusion by vein wall
 - To avoid hemolysis
 - If Microlance needle is used, collect sample by drip directly into specimen container.
- Release tourniquet.
- Remove needle and apply local pressure with dry gauze for 3 minutes or until complete hemostasis.



FIG. 13.5. A: Anatomy of the femoral triangle as defined in the text. (Adapted from Plaxico DT, Bucciarella RL. Greater saphenous vein venipuncture in the neonate. J Pediatr. 1978;93:1025, with permission) B: Position of the femoral triangle on the abducted thigh.

Scalp Vein

- Shave adequate area of frontal or parietal scalp.
- Use scalp vein needle set or 23-gauge butterfly.
- Occlude vein proximally with finger, or place a rubber band around head circumference, avoiding eye area.
- Feel for a pulse to avoid tapping an artery.
- Use a shallow angle (15 to 20 degrees).
- See technique for general venipuncture.

Proximal Greater Saphenous Vein (9)

- Have assistant hold infant's thighs abducted with knees and hips slightly flexed.
- Locate femoral triangle (Fig. 13.5A).
 - Proximal boundary: inguinal ligament
 - Lateral boundary: medial border of sartorius muscle
 - Medial boundary: lateral border of adductor longus muscle
- Enter skin and then vein at point approximately two-thirds along line from inguinal ligament to apex of triangle (Fig. 13.5B).
 - Use relatively steep angle (60 to 90 degrees).
 - After entering skin, advance while applying gentle suction 1 to 4 mm until blood return is achieved.
- See F, General Venipuncture.

External Jugular Vein

- Position infant in head-down position with head extended and rotated away from selected vessel (Fig. 13.6).
- Prepare skin over sternocleidomastoid muscle with antiseptic.
- Flick infant's heel to induce crying and optimize vein distension.
- Visualize external jugular vein running from angle of jaw to posterior border of sternocleidomastoid in its lower third.
- Puncture vessel where it runs across the anterior border of the sternocleidomastoid muscle.
- See F, "General Venipuncture.



FIG. 13.6. Infant positioned for puncture of external jugular vein.

- G. Complications (10,11,12 and 13)
 - Hemorrhage with
 - Coagulation defect
 - Puncture of deep vein
 - Venous thrombosis or embolus, with puncture of large, deep vein (11)
 - Laceration of adjacent artery
 - During femoral vein puncture:
 - Reflex arteriospasm of femoral artery with gangrene of extremity (12)
 - Penetration of peritoneal cavity
 - Septic arthritis of hip (13)
 - During internal jugular puncture:
 - Laceration of carotid artery
 - o Pneumothorax/subcutaneous emphysema
 - Interference with ventilation owing to positioning for jugular vein puncture
 - Raised intracranial pressure owing to head-down position aggravating intraventricular hemorrhage
 - During scalp vein puncture:
 - Laceration of artery

• Corneal abrasion or other eye damage if rubber band used improperly

References

1. Baral J. Use of a simple technique for the collection of blood from premature and full-term babies. Med J Aust. 1968;1:97.

2. Shohat M. Preterm blood counts vary with sampling site [Letter]. Arch Dis Child. 1987;62:1193.

3. Thurlbeck SM, McIntosh N. Preterm blood counts vary with sampling site. Arch Dis Child. 1987;62:72.

4. Shah VS, Taddio A, Bennett S, et al. Neonatal pain response to heel stick vs venipuncture for routine blood sampling. Arch Dis Child Fetal Neonat Ed. 1997;77:F143.

5. Larsson BA, Tannfeldt G, Lagercrantz H, et al. Alleviation of the pain of venipuncture in neonates. Acta Paediatr. 1998;87:774.

6. Archarya AB, Annamali S, Taub NA, Field D. Oral sucrose analgesia for preterm infant venipuncture. Arch Dis Child Fetal Neonat Ed. 2004;89:F17.

7. Stevens B, Yamada J, Ohlsson A. Sucrose for analgesia in newborn infants undergoing painful procedures. Cochrane Database Syst Rev. 2004;(3):CD001069.

8. Larsson BA, Tannfeldt G, Lagercrantz H, et al. Venipuncture is more effective and less painful than heel lancing for blood tests in neonates. Pediatrics. 1998;101:882

9. Plaxico DT, Bucciarelli RL. Greater saphenous vein venipuncture in the neonate. J Pediatr. 1978;93:1025. 10. McKay RJ Jr. Diagnosis and treatment: risk of obtaining samples of venous blood in infants. Pediatrics. 1966;38:906.

11. Nabseth DC, Jones JE. Gangrene of the lower extremities of infants after femoral venipuncture. N Engl J Med. 1963;268:1003.

12. Kantr RK, Gorton JM, Palmieri K, et al. Anatomy of femoral vessels in infants and guidelines for venous catheterizations. Pediatrics. 1989;33:1020.

13. Asnes RS, Arendar GM. Septic arthritis of the hip: a complication of venipuncture. Pediatrics. 1966;38:837.

14-Arterial Puncture

Dawn M. Walton Billie Lou Short

A. Indications (1, 2)

- Sampling for arterial blood gas determination
- Sampling for routine laboratory test when venous and capillary sampling not suitable or unobtainable

B. Contraindications

- Coagulation defects, thrombocytopenia
- Circulatory compromise in the extremity
- Inappropriate artery
 - Femoral artery
 - Use of radial artery if inadequate collaterals (see Allen's test below)
- Infection in sampling area
- When cannulation of that vessel is anticipated
- Use of peripheral arteries on the ipsilateral arm in an infant with congenital heart disease requiring a shunt via the subclavian artery

C. Precautions

- Perform arterial sampling only when venous or capillary sampling is inappropriate.
 - Arterial blood gas analyses prior to placement of indwelling access.
 - Ammonia levels
 - Large quantities of blood to be obtained
 - o Very low-birthweight infants with poor venous access
 - Use smallest possible needle to minimize trauma to vessel (23 to 27 gauge).
- Avoid laceration of the artery caused by puncturing both sides of arterial wall in exactly opposite locations.
- Guarantee hemostasis at end of procedure.
- Check distal circulation after puncture.
 - Arterial pulse
 - Capillary refill time
 - Color, temperature
- Take action to reverse arteriospasm, if necessary (see Chapter 33).

D. Selection of Arterial Site

- Peripheral site preferred
- Radial artery preferred if ulnar collateral intact
- Dorsalis pedis, posterior tibial arteries satisfactory
- Brachial artery only if urgent indication and no more peripheral arterial or umbilical artery access is available (3)
- Temporal artery should be avoided because of risk of neurologic damage (4, 5).

E. Equipment

- Gloves
- Needle
 - A 23- to 25-gauge venipuncture needle
 - Safety-engineered needle should be used
- Appropriate syringes
- Materials for minor skin preparation; povidone–iodine solution preparation preferred for blood culture
- Gauze pads
- Sterile glove to cover transilluminator
- High-intensity fiber-optic light for transillumination (see Chapter 12)
- Oral sucrose solution (24% to 25%) for pain control, if indicated (6, 7)

F. Technique

General Principles (1, 2)

- Transillumination may assist location of vessel (Fig. 12.1) (8).
- Position needle for arterial puncture against direction of blood flow.
 - Keep angle of entry shallow for superficial vessels.
 - 15 to 25 degrees for superficial artery, bevel down
 - 5 degrees for deep artery, bevel up
 - Penetrate skin first, and then puncture artery to minimize trauma to vessel.

 \circ Use fresh needle and repeat skin preparation if withdrawal from skin is necessary.

• Apply firm, local pressure for 5 minutes to achieve complete hemostasis.



FIG. 14.1. Position of wrist for puncture of radial artery. (1) distal wrist crease; (2) proximal wrist crease.

Radial Artery Puncture (2)

- Extend wrist, supine, not hyperextended (Fig. 14.1), which may occlude vessel.
 - Locate radial and ulnar arteries at proximal wrist crease (Fig. 14.2).
 - Radial artery is lateral to flexor carpi radialis tendon.
 - Ulnar artery is medial to flexor carpi ulnarius tendon.
 - Transillumination may be helpful.
- Perform modified Allen's test for collateral supply (9).
 - Elevate infant's hand.
 - Occlude both radial and ulnar arteries at wrist.
 - Massage palm toward wrist.
 - Release occlusion of ulnar artery only.
 - Look for color to return to hand in less than 10 seconds, indicating adequate collateral supply.
 - Do not puncture radial artery if color return takes more than 15 seconds.



FIG. 14.2. Anatomy of the major arteries of the wrist and hand.



В

А

FIG. 14.3. A: Penetration of artery at angle of 15 to 25 degrees with bevel down. Preferred method for small premature infants. B: Penetration of artery at angle of 45 degrees with bevel up.

- Locate artery by palpation and/or transillumination.
- Prepare area with antiseptic, as for minor procedure.
- Check function of syringe.
- Puncture skin and penetrate artery at 45 degrees with bevel up. For very small infants, use angle of 15 to 25 degrees with bevel down (Fig. 14.3).
 - While maintaining gentle suction, advance until there is blood return or resistance from bone.
 - If no blood obtained prior to encountering resistance, withdraw needle cautiously until blood returns. Artery may spasm when needle is introduced. Be patient; change angle of needle as necessary.
- Collect sample and remove needle.
- Compress site for 5 minutes or until hemostasis is complete.
- Verify satisfactory peripheral blood flow (10, 11).

Dorsalis Pedis Puncture

- Locate artery by palpation and transillumination on dorsum of foot between extensor hallucis longus and extensor digitorum longus tendons (Fig. 14.4).
- Choose an angle of 15 to 25 degrees.
- See F, General Principles.

Posterior Tibial Puncture

- Locate artery by palpation and transillumination between Achilles tendon and medial malleolus (Fig. 14.5).
- Choose an angle of 45 degrees.
- See F, "General Principles.



FIG. 14.4. Anatomic relations of the dorsalis pedis artery.





G. Complications (10)

See Chapter 30 for complications of arterial cannulation.

- Distal ischemia from arteriospasm, hematoma, thrombosis, or embolism
- Infection
 - Osteomyelitis (12)
 - Infected hip joint after femoral puncture (12)
- Hemorrhage or hematoma
- Nerve damage (13)
 - Median nerve (brachial artery puncture)
 - Posterior tibial nerve
 - Femoral nerve
- Extensor tendon sheath injury, resulting in "false cortical thumb†(14)
- Forearm compartment syndrome following brachial artery puncture (15)
- Inaccuracy of blood gas estimated (16, 17)
 - Excessive heparinization of syringe (falsely low PCO₂ and pH)
 - Hypothermic or hyperthermic infant
 - Gas bubbles in syringe
 - Spuriously high PO₂
 - Spuriously low PCO₂
 - Excessive delay in processing
 - Clot in syringe

References

1. Smith AD. Arterial blood sampling in neonates. Lancet. 1975;1:254.

2. Shaw JC. Arterial sampling from the radial artery in premature and full-term infants. Lancet. 1968;2:389.

3. Okeson GC, Wulbrecht PH. The safety of brachial artery puncture for arterial blood sampling. Chest. 1998;114:748.

4. Bull MJ, Schreiner RL, Garg BP, et al. Neurologic complications following temporal artery catheterization. J Pediatr. 1980;96:1071.

5. Simmons MA, Levine RL, Lubchenco LO, et al. Warning: serious sequelae of temporal artery catheterization. J Pediatr. 1978;92:284.

6. Acharya AB, Annamali S, Taub NA, Field D. Oral sucrose analgesia for preterm infant venipuncture. Arch Dis Childhood Fetal Neonatal Ed. 2004;89:F17.

7. Stevens B, Yamada J, Ohlsson A. Sucrose for analgesia in newborn infants undergoing painful procedures. Cochrane Database Syst Rev. 2004;(3):CD001069.

Wall PM, Kuhns LR. Percutaneous arterial sampling using transillumination. Pediatrics. 1977;59:1032.
 Cable DG, Mullany CJ, Schaff HV. The Allen test. Ann Thorac Surg. 1999;67:876.

10. Gillies ID, Morgan M, Sykes MK, et al. The nature and incidence of complications of peripheral artery puncture. Anaesthesia. 1979;34:506.

11. Noreng MF. Blood flow in the radial artery before and after arterial puncture. Acta Anaesthesiol Scand. 1986;30:281.

12. Nelson DL, Hable KA, Matsen JM. Proteus mirabilis osteomyelitis in two neonates following needle puncture. Successful treatment with ampicillin. Am J Dis Child. 1973;125:109.

13. Pape KE, Armstrong DL, Fitzhardinge PM. Peripheral median nerve damage secondary to brachial arterial blood gas sampling. J Pediatr. 1978;93:852.

14. Skogland RR, Giles EJ. The false cortical thumb. Am J Dis Child. 1986;140:375.

15. Safran MR, Bernstein A, Lesavoy MA. Forearm compartment syndrome following brachial arterial puncture in uremia. Ann Plast Surg. 1994;32:535.

16. Fan LL, Dellinger KT, Mills AL. Potential errors in neonatal blood gas measurements. J Pediatr. 1980;97:650.

17. Goldsmith JP. Error in blood gas sampling results in spurious interpretation of compensated metabolic acidosis. J Perinatol. 1993;13:165.

15-Capillary Heelstick Blood Sampling

Laura A. Folk

A. Definitions

- Automated heel lancing device: Encased, spring-loaded, retractable blade that provides a controlled and consistent width and depth of incision for blood testing.
 - Incision depths range from 0.65 to 2.0 mm for micropreemies through toddlers (Tenderfoot, International Technidyne Corporation, Edison, NJ, USA) and from 0.85 to 1.0 mm for preemies and newborns (BD Quickheel Lancet, BD Vacutainer Systems, Franklin Lakes, NJ, USA) (Table 15.1).
 - The controlled depths avoid damage to the calcaneus (1,2) while providing greater yield with less pain, hemolysis, and laboratory-value error (3,4,5 and 6). The shallower devices can be used to obtain small samples from larger infants who require frequent point-of-care glucose testing (7).
 - Nonautomated (manual) stylet-type lancets and spring-loaded needle-puncture devices designed for adult glucose testing are not appropriate for infants (6).
- Heelwarmer: Chemically activated packet to heat heel prior to capillary testing. If heel warming is used, a commercial prepackaged unit provides controlled temperature. The warmer should be applied for 5 minutes and then removed prior to heelstick.

B. Purpose

To obtain capillary blood samples that provide accurate laboratory results with minimal discomfort and potential for injury/infection.

C. Background

Capillary heel sampling is an easily mastered, minimally invasive technique that, when performed with proper technique and equipment, provides laboratory results within acceptable tolerances compared with samples from arterial catheters (8). Capillary blood sampling conserves veins in premature and critically ill infants, who can be expected to require intravenous access for fluids or medication administration.

D. Indications

- Capillary blood gas sampling
- Routine laboratory analysis (standard hematology, chemistries, toxicology/drug levels) requiring a limited amount of blood in which minimal cell lysis does not alter results
- Newborn metabolic screen

E. Contraindications

- Edema, because interstitial fluid dilutes the sample and gives inaccurate results
- Injury or anomalies that preclude putting pressure on the foot
- Areas that are bruised or injured by multiple previous heelsticks
- Poor perfusion
- Local infection

F. Limitations

- Venous or arterial blood rather than capillary samples should be used for:
 - Blood cultures, which require sterile technique
 - Tests in which even a minimal amount of hemolysis will compromise results
 - Special tests such as coagulation studies (newer coagulation tests that require only a few drops of blood are still not widely available)
 - Laboratory tests that require more than 1.5 mL of blood

G. Equipment

- Gloves
- Heel-warming device if desired
- Antiseptic (betadine/saline or alcohol swab)
- Pad or other means of protecting bed linens
- Heel-lancing device. Use appropriate size for infant (Table 15.1).
- Specimen collector as appropriate
 - Serum separators
 - Hematology tubes
 - Capillary blood gas tube
 - Newborn metabolic screen filter paper
- Capillary tubes for blood transfer to lab tubes if appropriate
- Bandage or gauze wrap

TABLE 15.1 Examples of Automated free Earcing Froducts Dased on Infant Size		
Infant Size	Available Products	Incision Depth/Length
< 1,000 g	Tenderfoot Micro-preemie	0.65 mm/1.40 mm
Low-birthweight & preemie <1,000 g	gTenderfoot Preemie/BD Quickheel Preemie	e0.85 mm/1.75 mm
Term to 3-6 mo	Tenderfoot Newborn/BD Quickheel Infant	1.0 mm/2.5 mm
6 mo-2 years	Tenderfoot Toddler	2.0 mm/3.00 mm

TABLE 15.1 Examples of Automated Heel Lancing Products Based on Infant Size

H. Precautions

- Site
 - Do not use the end of the heel. The calcaneum is superficial at this site, and there is an 0 increased risk of osteomyelitis (1).
 - Do not use fingertips, toes, or earlobes of babies.
- Hand position •
 - Do not squeeze the heel. Squeezing the heel results in greater pain, lower blood yield, and increased cell lysis.
- Collection •
 - If using capillary tubes for blood transfer, it is essential to determine whether the tube contains substances such as anticoagulants, and whether those substances have the potential to interfere with lab results. Do not use tubes containing anticoagulants for newborn metabolic screens.
 - Scoop-shaped collectors provided with mini–lab tubes are used to guide blood drops to the 0 specimen tube. Avoid repeated "scooping†along the surface of the foot. Microclots that form in blood on the skin can alter lab results.



FIG. 15.1. Appropriate sites for capillary heelstick sampling are along the sides of the heels.



FIG. 15.2. Alternative site for capillary heelstick sampling. If frequent sampling has rendered the sides of

the heels unsuitable, the plantar surface between them can be used. Do not incise the end of the heel.

I. Technique

- Identify site; the preferred areas for capillary heel testing are the outer aspects of the heel (Fig. 15.1).
 - Vary sites to prevent bruising and skin damage.
 - The plantar surface can be used if the preferred areas are compromised by previous frequent testing (Fig. 15.2) (1,8). The skin-to-calcaneal perichondrium distance is at least 3 mm in most term babies and in 91% of babies at 33 to 37 weeks' gestation, but is at least 3 mm in only about 60% of babies <33 weeks' gestation (1).
- Apply heel warmer or warm towel for 5 minutes. Remove just before procedure.
- Provide comfort measuresâ€"Facilitated tucking/swaddling and the use of pacifiers combined with administration of a concentrated sucrose solution results in less measurable pain and faster resolution of discomfort in the infant following the procedure (9,10) (Fig. 2.1).
- Prepare automated device by removing release clip.
- Don gloves.
- Cleanse site with betadine followed with saline wipe or alcohol wipe.
- Position hand with fingers along the calf and thumb at ball of foot to stabilize. Apply pressure along calf toward heel (Fig. 15.3).
- Place automated device on site and activate.
- Apply pressure to leg with counterpressure to ball of foot and allow blood drop to form.
- Wipe away first drop of blood with gauze or clean wipe.
- Using capillary action, fill blood gas tube, holding tube horizontally (Fig. 15.4).
- Release pressure, allowing capillaries to refill.
- Guide blood drops into tube or collect with capillary tube for transfer to laboratory tube.
- If blood stops flowing, wipe site to remove clot with alcohol swab, gauze, or clean wipe, ensure time for capillary refill, and then reapply pressure to leg. If blood does not flow, choose another site and repeat procedure or consider venipuncture.
- When samples have been collected, apply pressure to puncture site and wrap with gauze or apply adhesive bandage.
- Continue comfort measures.



FIG. 15.3. Position for hand and automated lancing device. Position heel in the apex of the angle of the thumb and forefinger with fingers along the calf and thumb along the ball of the foot. Position automated lancing device in appropriate position. Apply pressure along the calf with counterpressure by the thumb. Do not squeeze the heel.



FIG. 15.4. Capillary blood gas sampling.

J. Specimen Handling

- Collect blood gas sample first, then hematology samples, then chemistry/toxicology samples.
- Ensure that blood gas samples are free of air bubbles.
 - Place the tube horizontally so that the blood is drawn by capillary action and does not collect air bubbles that can alter results. Apply caps to ends of tube.
 - Capillary blood gas samples should be analyzed within 10 minutes or should be kept horizontally on ice for up to 1 hour, and the tube must be rolled prior to analysis. Consult institution laboratory for guidance on blood gas sample storage and transport.
- Flick side of hematology microtube during collection process to activate anticoagulant and prevent clotting.
- Newborn metabolic screen: specific collection guidelines (11)
 - Minimum 24 to 48 hours after birth
 - Integrity of collection medium: Avoid touching filter paper, as oils from finger can compromise results.
 - Single (no overlapping) drops on filter paper. Position infant so that incision is down, allowing a large drop of blood to form. Blood should drop freely onto designated circle on filter paper. Repeat for each circle.
 - Do not apply blood using capillary tubes that contain anticoagulants or other materials that can interfere with lab results.



FIG. 15.5. Cellulitis of heelâ€" complication of capillary heelstick sampling.

- K. Complications
 - Pain
 - \circ $\;$ Provide oral sucrose, swaddling/tucking, and pacifier.
 - Use proper equipment.
 - Make incisions in designated areas of the heel.
 - Infection (cellulitis, abscess, perichondritis, osteomyelitis) (Fig. 15.5) (12,13)
 - Tissue loss and scarring
 - Calcified nodules (14)

L. Inaccurate laboratory results

- Hyperkalemia secondary to excessive hemolysis
 - Use proper technique and procedures to minimize cell lysis.
- Erroneous blood gas results
 - Ensure that sample is free of air bubbles.
 - Avoid delay in analysis.
 - Use proper technique and procedures to minimize cell lysis.

References

1. Arena J, Emparanza J, Nogues A, Burls A. Skin to calcaneus distance in the neonate. Arch Dis Child Fetal Neonatal Ed. 2005;90(4):F328–F331.

2. Vertanen H, Fellman V, Brommels M, Viinikka L. An automated incision device for obtaining blood samples from the heels of preterm infants causes less damage than a conventional manual lancet. Arch Dis Child Fetal Neonatal Ed. 2001;84(1):F53–F55.

3. Shepherd AJ, Glenesk A, Niven CA, Mackenzie J. A Scottish study of heel-prick blood sampling in newborn babies. Midwifery. 2005;7(2): 158–168.

4. Kellan B, Waller J, McLaurin C, et al. Tenderfoot preemie vs a manual lancet: a clinical evaluation. Neonatal Network. 2001;20(7):31–36.

5. Kazmierczak SC, Robertson AF, Briley KP. Comparison of hemolysis in blood samples collected using an automatic incision device and a manual lancet. Arch Pediatr Adolesc Med. 2002;156(11):1072–1074.
6. Shah V, Taddio A, Kulasekaran K, et al. Evaluation of a new lancet device (BD QuickHeel) on pain response and success of procedure in term neonates. Arch Pediatr Adolesc Med. 2003;157(11):1075–1078.

7. Noerr B. State of the science: neonatal hypoglycemia. Adv Neonatal Care. 2001;1(1):4–21.

Solution K, Cress G, Connolly N, et al. Neonatal laboratory blood sampling: comparison of results from arterial catheters with those from an automated capillary device. Neonatal Network. 2000;19(1):27–34.
 Gibbins S, Stevens B. The influence of gestational age on the efficacy and short-term safety of sucrose for

procedural pain relief. Adv Neonatal Care. 2003;3(5):241–249. 10. Coleman M, Solarin K, Smith C. Assessment and management of pain and distress in the neonate. Adv Neonatal Care. 2002;2(3):123–139.

11. Bryant K, Horns K, Longo N, Schiefelbein J. A primer on newborn screening. Adv Neonatal Care. 2004;4(5):306–317.

12. Abril Martin JC, Aguilar Rodriguez L, Albinana Cilvetti J. Flatfoot and calcaneal deformity secondary to osteomyelitis after heel puncture. J Pediatr Orthop. 1999;8:122–124.

13. Lauer BA, Altenburgher KM. Outbreak of staphylococcal infections following heel puncture for blood sampling. Am J Dis Child. 1981;135:277–278.

14. Williamson D, Holt PJ. Calcified cutaneous nodules on the heels of children: a complication of heelsticks as a neonate. Pediatr Dermatol. 2001;18:138–140.

16-Lumbar Puncture

S. Lee Woods

A. Indications (1,2)

• To diagnose central nervous system (CNS) infections (meningitis, encephalitis), including congenital infections (TORCHâ€"toxoplasmosis, other infections [usually implying syphilis], rubella, cytomegalovirus, and herpes simplexâ€"infections) as well as bacterial and fungal infections

Routine inclusion of lumbar puncture (LP) in the initial sepsis evaluation of newborn infants (in the first 7 days of life) is controversial (3,4,5,67 and 8). Meningitis occurs less frequently in this population than in older newborns, and the majority of cases of meningitis occur in infants with positive blood cultures. The procedure may be poorly tolerated by newborns with cardiorespiratory compromise (910,11). An LP is indicated if early bacteremia is documented or if signs of CNS involvement are present (seizures, coma, focal neurologic abnormality). An LP is also indicated in the evaluation for acquired infection in the later neonatal period, when the incidence of meningitis is significant. In a recent review (8), as many as one third of very low-birthweight infants who had lateonset meningitis (after 3 days of life) did so in the absence of positive blood culture.

- To monitor efficacy of antimicrobial therapy in the presence of CNS infection by examining cerebrospinal fluid (CSF) cell count, microbiology, and drug levels (12)
- To drain CSF in communicating hydrocephalus associated with intraventricular hemorrhage (1,2,13)

For effective treatment of posthemorrhagic hydrocephalus by this means, there must be communication between the lateral ventricles and the lumbar subarachnoid space, and an adequate volume of CSF (10 to 15 mL/kg) must be obtained. Communication is demonstrated by an immediate decrease in ventricular size or change in anterior fontanelle or head circumference following LP. Efficacy and safety of serial LPs in the temporary amelioration or long-term improvement of posthemorrhagic hydrocephalus are controversial (14,15,16,17 and 18). Potential risks of repeated LPs must be weighed against possible benefits.

- To aid in the diagnosis of metabolic disease (1,2,19)
- To diagnose intracranial hemorrhage

The finding of increased red blood cells and protein content in the CSF or xanthochromia of centrifuged fluid suggests intracranial hemorrhage. The definitive diagnosis and determination of the site of hemorrhage (subdural, subarachnoid, intraparenchymal, intraventricular) are best made by neuroimaging techniques such as computed tomography (CT) or magnetic resonance imaging (MRI).

- To diagnose CNS involvement with leukemia
- To inject chemotherapeutic agents
- To instill contrast material for myelography

B. Contraindications (1,2,20,21)

• Increased intracranial pressure (ICP)

Increased ICP may occur with bacterial meningitis or intracranial mass lesions. In the neonate with open cranial sutures, this rarely results in signs of transtentorial or cerebellar herniation. However, herniation can occur after LP in the presence of elevated ICP, even when the sutures are open. If

signs of significant increased ICP exist (rapidly declining or severely depressed level of consciousness, abnormal posturing, cranial nerve palsies, tense anterior fontanelle, abnormalities in heart rate, respirations, or blood pressure without other cause), CT or MRI should be performed before LP. Papilledema is a late sign and is rarely present in the neonate, regardless of the degree of increased ICP.

- Uncorrected thrombocytopenia or bleeding diathesis
- Infection in the skin or underlying tissue at or near the puncture site
- Lumbosacral anomalies
- Cardiorespiratory instability, which may be exacerbated by the procedure

C. Equipment

All equipment must be sterile, except facemask. Prepackaged lumbar puncture kits are available.

- Gloves and mask
- Cup with iodophor antiseptic solution
- Gauze swabs
- Towels or transparent aperture drape
- Spinal needle with short bevel and stylet, 20 or 22 gauge x $1\frac{1}{2}$ in
- Three or more specimen tubes with caps
- Adhesive bandage

D. Precautions

- Monitor vital signs and oxygen saturation. Preoxygenation and increased supplemental oxygen during the procedure can prevent hypoxemia (22). Airway compromise can be reduced by avoiding the fully flexed lateral decubitus position and direct flexion of the neck (9,10 and 11). Flexing the hips to only 90 degrees avoids abdominal compression and the potential for aspiration.
- Use strict aseptic technique as for a major procedure (see Chapter 4).
- Always use a needle with stylet to avoid development of intraspinal epidermoid tumor (23).

Incidence of traumatic LP is not reduced by use of a needle without stylet (24).

- To prevent traumatic tap caused by overpenetration, insert the needle slowly while removing the stylet at frequent intervals to detect CSF as soon as the subdural space is entered.
- Never aspirate CSF with a syringe. Even a small amount of negative pressure can increase the risk of subdural hemorrhage or herniation.
- Palpate landmarks accurately to prevent puncture above the L2-L3 interspace (lower interspace should be used for preterm infants; see discussion under E.2).



FIG. 16.1. Restraining infant for lumbar puncture in the lateral recumbent position. Neck should not be flexed.

E. Technique (20,21,25,26 and 27)

- Have an assistant restrain the infant in the lateral decubitus or sitting position, with spine flexed (Figs. 16.1 and 16.2). Avoid flexion of the neck, as this increases the chance of airway compromise.
- Palpate the interspace that falls immediately above or below an imaginary line drawn between the iliac crests (L3–L4 and L4–L5 interspaces, preferred sites for LP) (Fig. 16.3).

The termination of the spinal cord relative to the spine changes during fetal development and early infancy (28,29). The normal adult termination, between the middle of T12 and the lower portion of L3 vertebrae (28,30), is not achieved until 2 months postterm (28). Between 25 and 40 weeks' gestation, the cord termination gradually ascends from L4 to L2 (28). This should be taken into account and the lower L4–L5 interspace used for lumbar puncture in significantly preterm infants to avoid possible cord penetration (29).

- Prepare as for major procedure (see Chapter 4). Wash hands thoroughly. Put on mask and sterile gloves.
- Clean the lumbar area three times with antiseptic.
 - Begin at the desired interspace and wash in enlarging circles to include the iliac crests.
 - Allow antiseptic to dry or blot excess with sterile gauze.
- Drape, leaving the puncture site and infant's face exposed. A transparent aperture drape is recommended because it does not obstruct the view of the patient.

Local anesthesia is generally not used for LP in neonates. Use of local anesthetic cream prior to cleaning the area may be helpful in reducing pain during LP (31,32 and 33). Use of lidocaine injection does not reduce physiologic instability but may reduce struggling by the infant during the procedure (34,35).

- Insert the needle in the midline into the desired interspace.
 - Aim slightly cephalad (on a plane with the umbilicus) to avoid the vertebral bodies (Fig. 16.4).

- If resistance is met, withdraw the needle slightly and redirect more cephalad.
- Hold a finger on the vertebral process above the interspace to aid in locating the puncture site if the infant moves.



FIG. 16.2. Restraining infant for lumbar puncture in the sitting position.



FIG. 16.3. A: Externally palpable anatomic landmarks. B: Vertebral bodies removed to show anatomy of spinal cord in lumbosacral area in relation to external landmarks.

- Advance the needle slowly to a depth of approximately 1 to 1.5 cm in a term infant, less in a preterm infant, until the epidermis and dermis are traversed.
 - As the needle is further advanced, remove the stylet frequently to check for fluid. Replace the stylet before advancing the needle.
 - A change in resistance can often be felt as the needle passes through the ligamentum flavum and dura (Fig. 16.5). This may be more difficult to appreciate in a young infant than in an older child.
 - Wait for fluid after removing the stylet, as the flow may be slow.
 - If no fluid is obtained, rotate the needle to reorient the bevel. If no fluid is obtained, replace the stylet, remove the needle, and try one interspace above or below, using a new needle for each attempt.
- Collect CSF for diagnostic studies. Allow CSF to flow passively into the collection tubes; never aspirate with a syringe. Accurate opening pressure measurement is possible in a quiet infant.
 - Collect 1 mL of CSF in each of three to four tubes.
 - Send first sample for bacterial culture.
 - Send last sample for cell count, unless fluid becomes visibly more bloody during the tap.
 - Send the remainder for desired chemical and microbiologic studies.
 - Look for clearing of fluid in successive collections in the event of a traumatic tap.
- For myelography or instillation of chemotherapeutic agents, it is not necessary to remove CSF.
- For treatment of hydrocephalus, remove 10 to 15 mL/kg of CSF, or collect until CSF flow ceases (up to 10 minutes).
- Replace the stylet before removing the needle to prevent entrapment of spinal nerve roots in the extradural space. Remove the needle, and place an adhesive bandage over the puncture site.


FIG. 16.4. Inserting spinal needle in slightly cephalad direction to avoid vertebral bodies.



FIG. 16.5. Needle has penetrated the dura, and stylet has been removed to allow free flow of spinal fluid.

F. Complications (1,2,36)

In older children and adults, headache is the most common complication following LP, occurring in up to 40% of patients (36). There is no clear evidence that headache occurs in infants. In neonates, the most common complication is transient hypoxemia from positioning for the procedure (9,10 and 11). In some reports, this is seen in a majority of cases and depends on the method of positioning used. Also common, occurring in up to 36% of LPs in neonates (27), is contamination of the CSF sample with blood from puncture of the epidural venous plexus on the posterior surface of the vertebral body (traumatic tap). All other potential complications listed below are rare, occurring in about 0.3% of LPs (36).

- Hypoxemia from kneeâ€"chest position (9,10 and 11)
- Contamination of CSF sample with blood (traumatic tap) (27)
- Aspiration
- Cardiopulmonary arrest
- Sudden intracranial decompression with cerebral herniation (37,38)
- Infection
 - Meningitis from LP performed during bacteremia (incidence about 0.2%) (39,40 and 41)
 - o Discitis (42)
 - Spinal cord abscess (43)
 - Epidural abscess (43,44)
 - Vertebral osteomyelitis (44)

- Bleeding
 - Spinal epidural hematoma (45)
 - Spinal or intracranial subdural hematoma (46,47 and 48)
 - Spinal or intracranial subarachnoid hematoma (48,49)
 - Rupture of intracranial aneurysm (48)
- Intraspinal epidermoid tumor from epithelial tissue introduced into the spinal canal (23)
- Spinal cord puncture and nerve damage if puncture site is above the level of cord termination (see discussion in E.2 concerning cord termination in preterm infants) (29)
- Sixth-nerve palsy caused by removal of excessive CSF with resulting traction on the nerve (50)
- Deformity of the lumbar spine secondary to acute spondylitis (51)

References

1. Michelson DJ. Spinal fluid evaluation. In: Swaiman KF, Ashwal S, Ferriero DM, eds. Pediatric Neurology. 4th ed. Philadelphia: Mosby; 2006:153.

2. Menkes JH, Moser FG. Neurologic examination of the child and infant. In: Menkes JH, Sarnat HB, Maria BL, eds. Child Neurology. 7th ed. Philadelphia: Lippincott Williams & Wilkins; 2006:1.

3. Albanyan EA, Baker CJ. Is lumbar puncture necessary to exclude meningitis in neonates and young infants: lessons from the group B streptococcus cellulitis–adenitis syndrome. Pediatrics. 1998;102:984.

4. Johnson CE, Whitwell JK, Pethe K, et al. Term newborns who are at risk for sepsis: are lumbar punctures necessary? Pediatrics. 1997;99:e10.

5. Beerman MR, Chopde N, Dawood Y, et al. Lumbar puncture in the evaluation of possible asymptomatic congenital syphilis in neonates. J Pediatr. 1996;128:125.

6. Wiswell TE, Baumgart S, Gannon CM, et al. No lumbar puncture in the evaluation for early sepsis: will meningitis be missed? Pediatrics. 1995;95:803.

7. Kumar P, Sarkar S, Narang A. Role of routine lumbar puncture in neonatal sepsis. J Paediatr Child Health. 1995;31:1.

8. Stoll BJ, Hansen N, Fanaroff AA, et al. To tap or not to tap: high likelihood of meningitis without sepsis among very low birth weight infants. Pediatrics. 2004;113:1181.

9. Weisman LE, Merenstein GB, Steenbarger JR. The effect of lumbar puncture position in sick neonates. Am J Dis Child. 1983;137:1077.

10. Gleason CA, Martin RJ, Anderson JV, et al. Optimal position for a spinal tap in preterm infants. Pediatrics. 1983;71:31.

11. Spahr RC, MacDonald HM, Mueller-Heubach E. Knee–chest position and neonatal oxygenation and blood pressure. Am J Dis Child. 1981;135:79.

12. Wubbel L, McCracken GH. Management of bacterial meningitis 1998. Pediatr Rev. 1998;19:78.

13. Kreusser KL, Tarby TJ, Kovnar E, et al. Serial lumbar punctures for at least temporary amelioration of neonatal posthemorrhagic hydrocephalus. Pediatrics. 1985;75:719.

14. Ventriculomegaly Trial Group. Randomized trial of early tapping in neonatal posthaemorrhagic ventricular dilatation: results at 30 months. Arch Dis Child. 1994;70:F129.

15. Ventriculomegaly Trial Group. Randomized trial of early tapping in neonatal posthaemorrhagic ventricular dilatation. Arch Dis Child. 1990;65:3.

16. Dykes FD, Dunbar B, Lazarra A, et al. Posthemorrhagic hydrocephalus in high-risk preterm infants: natural history, management, and long-term outcome. J Pediatr. 1989;114:611.

17. Smith KM, Deddish RB, Ogata ES. Meningitis associated with serial lumbar punctures and posthemorrhagic hydrocephalus. J Pediatr. 1986;109:1057.

18. Bergman I, Wald ER, Meyer JD, et al. Epidural abscess and vertebral osteomyelitis following serial lumbar punctures. Pediatrics. 1983;72:476.

19. Hoffman GF, Surtees RA, Wevers RA. Cerebrospinal fluid investigations for neurometabolic disorders. Neuropediatrics. 1998;29:59.

20. Barone MA. Pediatric procedures. In: McMillan JA, Feigin RD, DeAngelis CD, et al., eds. Oski's Pediatrics: Principles and Practice. 4th ed. Philadelphia: Lippincott Williams & Wilkins; 2006:2671.

21. Haslam RHA. Neurologic evaluation. In: Behrman RE, Kliegman RM, Jenson HB, eds. Nelson Textbook of Pediatrics. 17th ed. Philadelphia: Saunders; 2004:1973.

22. Fiser DH, Gober GA, Smith CE, et al. Prevention of hypoxemia during lumbar puncture in infancy with preoxygenation. Pediatr Emerg Care. 1993;9:81.

23. Potgieter S, Dimin S, Lagae L, et al. Epidermoid tumours associated with lumbar punctures performed in early neonatal life. Dev Med Child Neurol. 1998;40:266.

24. Schreiner RL, Kleiman MB. Incidence and effect of traumatic lumbar puncture in the neonate. Dev Med Child Neurol. 1979;21:483.

25. Shilkofski N. Procedures. In: Robertson J, Shilkofski N, eds. The Harriet Lane Handbook. 17th ed. St. Louis, MO: Mosby; 2005:73.

26. Ferriero D, Buescher ES. Central nervous system. In: Taeusch HW, Christiansen RO, Buescher ES, eds. Pediatric and Neonatal Tests and Procedures. Philadelphia: Saunders; 1996:409.

27. Bonadio WA. Interpreting the traumatic lumbar puncture. Contemp Pediatr. 1989;6:109.

28. Barson AJ. The vertebral level of termination of the spinal cord during normal and abnormal development. J Anat. 1970; 106:489.

29. Tubbs RS, Smyth MD, Wellons JC, et al. Intramedullary hemorrhage in a neonate after lumbar puncture resulting in paraplegia. Pediatrics. 2004;113:1403.

30. Wilson DA, Prince JR. MR imaging determination of the location of the normal conus medullaris throughout childhood. Am J Roentgenol. 1989;152:1029.

31. Kaur G, Gupta P, Kumar A. A randomized trial of eutectic mixture of local anesthetics during lumbar puncture in newborns. Arch Pediatr Adolesc Med. 2003;157:1065.

32. Dutta S. Use of eutectic mixture of local anesthetics in children. Indian J Pediatr. 1999;66:707.

33. Taddio A, Ohlsson A, Einarson TR, et al. A systematic review of lidocaine–prilocaine (EMLA) in the treatment of acute pain in neonates. Pediatrics. 1998;101:e1.

34. Pinheiro JMB, Furdon S, Ochoa LF. Role of local anesthesia during lumbar puncture in neonates. Pediatrics. 1993;91:379.

35. Porter FL, Miller JP, Cole FS, et al. A controlled clinical trial of local anesthesia for lumbar punctures in newborns. Pediatrics. 1991;88:663.

36. Evans RW. Complications of lumbar puncture. Neurol Clin North Am. 1998;16:83.

37. Shetty AK, Desselle BC, Craver RD, et al. Fatal cerebral herniation after lumbar puncture in a patient with a normal computed tomography scan. Pediatrics 1999;103:1284.

38. Rennick G, Shann F, de Campo J. Cerebral herniation during bacterial meningitis in children. Br Med J 1993; 306:953.

39. Domingo P, Mancebo J, Blanch L, et al. Iatrogenic streptococcal meningitis. Clin Infect Dis. 1994;19:356.

40. Hristeva L, Booy R, Bowler I, et al. Prospective surveillance of neonatal meningitis. Arch Dis Child. 1993;69:14.

41. Eng RHK, Seligman SJ. Lumbar puncture-induced meningitis. JAMA. 1981;245:1456.

42. Bhatoe HS, Gill HS, Kumar N, et al. Post lumbar puncture discitis and vertebral collapse. Postgrad Med J. 1994;70: 882.

43. Bertol V, Ara JR, Oliveros A, et al. Neurologic complications of lumbar spinal anesthesia: spinal and paraspinal abscess. Neurology. 1997;48:1732.

44. Bergman I, Wald ER, Meyer JD, et al. Epidural abscess and vertebral osteomyelitis following serial lumbar punctures. Pediatrics. 1983;72:476.

45. Dulac O, Aicardi J, Lepintre J, et al. Hematome epidural intrarachidien apres ponction lombaire. Arch Fr Pediatr. 1975; 32:77.

46. Vos PE, De Boer WA, Wurzer JAL, et al. Subdural hematoma after lumbar puncture: two case reports and review of the literature. Clin Neurol Neurosurg. 1991;93:127.

47. Spanu G, Berlanda P, Rodriguez Y, et al. Spinal subdural haematoma: a rare complication of lumbar puncture. Neurochirurgia. 1988;31:157.

48. Hart IK, Bone I, Hadley DM. Development of neurological problems after lumbar puncture. Br Med J. 1988;296:51.

49. Blade J, Gaston F, Montserrat E, et al. Spinal subarachnoid hematoma after lumbar puncture causing reversible paraplegia in acute leukemia. J Neurosurg. 1983;58:438.

50. Bryce-Smith R, Macintosh RR. Sixth-nerve palsy after lumbar puncture and spinal analgesia. Br Med J. 1951;1:275.

51. Lintermans JP, Seyhnaeue V. Spondolytic deformity of the lumbar spine and previous lumbar punctures. Pediatr Radiol. 1977;5:181.

17-Subdural Tap

S. Lee Woods

- A. Indications (1,2,34)
 - To diagnose acute subdural collection over the cerebral convexities (hemorrhage, effusion, empyema) (5,6,7,8 and 9)

Computed tomography (CT), now generally available, is a safer method for detecting subdural fluid. Subdural tap should be reserved as a diagnostic tool for the infant who is too unstable to be transported for CT scanning.

- To sample convexity subdural collection for hematologic, microbiologic, and biochemical studies
- To drain convexity subdural collection to reduce increased intracranial pressure or to prevent the development of craniocerebral disproportion

Repeated therapeutic subdural taps should not be performed unless the infant is symptomatic or the head size is growing rapidly. Surgical intervention is indicated if subdural taps are not effective in controlling these symptoms (3).

B. Contraindications

- Clinical instability when risk exceeds potential benefit
- Uncorrected thrombocytopenia or bleeding diathesis
- Infection in the skin or underlying tissue at or near the puncture site

C. Equipment

All equipment must be sterile, except safety razor and facemask.

- Gloves and facemask
- Cup with iodophor antiseptic solution
- Gauze swabs
- Drapes or surgical towels
- Two short bevel needles, 19 to 22 gauge x 1 in, with stylets
- Specimen tubes with caps
- Adhesive bandage
- Safety razor

D. Precautions

• Use strict aseptic technique as for a major procedure (see Chapter 4).

- Insert the needle as far laterally as possible at the border of the anterior fontanelle or along the coronal suture, at least 1 to 2 cm from the midline, to avoid puncturing the sagittal sinus. Do not direct the needle medially during insertion.
- Remove the needle if there is not a definite change in resistance on penetrating the dura after insertion to approximately 0.5 to 1 cm.
- Hold the needle securely at all times to avoid inadvertent movement of the needle tip. Grasp the needle firmly or apply a hemostat at approximately 1 cm from the beveled end of the needle to prevent inadvertent advancement of the needle into the cerebral cortex.
- Allow fluid to drain spontaneously. Do not aspirate with a syringe.
- Limit fluid collected to 15 to 20 mL from each side. Removal of larger volumes can lead to bleeding into the subdural space.
- If frequent taps are required, vary the puncture site slightly to prevent fistula formation.
- Following the procedure, apply pressure to the scalp for 2 to 3 minutes to prevent fluid leak from the puncture site or subgaleal fluid collection.

E. Technique (1,10,11)

- Place the infant supine, with the crown of the head at the table edge. Monitor cardiorespiratory status.
- Have the assistant restrain the infant and steady the infant's head (Fig. 17.1).
- Shave the head over a wide area surrounding the anterior fontanelle (Fig. 17.1).
- Locate the junctions of the coronal sutures and anterior fontanelle.
- Put on mask. Wash hands thoroughly and put on sterile gloves.



FIG. 17.1. Position and restraint for subdural tap. Stippling demonstrates area to be prepared for procedure. An arrow indicates site for needle puncture.

- Clean the fontanelle and surrounding area three times with antiseptic solution. (See Chapter 4 for aseptic preparation for major procedure.)
 - Begin at the fontanelle and wash in enlarging circles.
 - Allow antiseptic to dry. Blot excess with sterile gauze.
- Cover infant's head with sterile drapes, leaving the anterior fontanelle and the infant's nose and mouth exposed.

• Locate the coronal suture by palpation at the lateral corner of the anterior fontanelle.

Generally, anesthesia is not required, but local injection of lidocaine at this time or application of topical anesthetic cream prior to cleaning the area can be used for local anesthesia at the puncture sites (1,12,13,14 and 15).

- Insert the needle slowly through the coronal suture, just lateral to its junction with the anterior fontanelle (see Fig. 17.1).
 - Hold the needle perpendicular to the skin surface.
 - Grasp the needle shaft with thumb and index finger, bracing the hand against the infant's head to maintain control of the needle during insertion (Fig. 17.2).
 - As the needle advances through the skin, pull the scalp slightly to create a Z-like track through the underlying tissue. This will help prevent fluid leakage from the puncture site or into the subgaleal space after removal of the needle.
- Advance until a "pop†is felt upon penetrating the dura. Remove the stylet (Fig. 17.2).
- Allow fluid to drain spontaneously into the sterile tubes until flow ceases or a maximum volume of 15 to 20 mL is reached. Fluid is sent for protein content, cell count, and culture.
- If no fluid appears, replace the stylet and remove the needle slowly. Do not reinsert on the same side.
- Repeat the procedure on the opposite side with a new, sterile needle.
- After removing the needle, apply firm pressure to the puncture site with sterile gauze for 2 to 3 minutes.
- Dress the puncture site with a small adhesive bandage.



FIG. 17.2. Coronal section of anatomic drawing showing subdural needle penetrating the dura in a patient with bilateral convexity subdural fluid collections. Operator's fingers are placed for maximal stabilization of the needle.

- F. Complications (10,11)
 - Subdural bleeding from laceration of the superior sagittal sinus or smaller vessels or from removal of excessive fluid with shift of intracranial contents and rebleeding
 - Development of chronic subdural fluid collection (16,17)

This complication may develop more frequently in infants treated with subdural tap. In one series of cases, 41.6% of patients treated with subdural tap developed chronic subdural collections, compared with 13% of those treated with craniotomy (17).

- Subgaleal fluid or blood accumulation
- Failure of the procedure to remove clotted subdural blood
- Infection

In one small series of cases, 1 of 12 infants (8%) treated with subdural tap developed subdural empyema after multiple taps (17).

- Trauma to the underlying cortex caused by inserting the needle too far
- Fistula formation after repeated taps

References

1. Haslam RHA. Neurologic evaluation. In: Behrman RE, Kliegman RM, Jenson HB, eds. Nelson Textbook of Pediatrics. 17th ed. Philadelphia: Saunders; 2004:1973.

2. Brett EM, Harding BN. Intracranial and spinal cord tumours. In: Brett EM, ed. Paediatric Neurology. 3rd ed. New York: Churchill Livingstone; 1997:537.

3. Volpe JJ. Intracranial hemorrhage: subdural, primary subarachnoid, intracerebellar, intraventricular (term infant), and miscellaneous. In: Neurology of the Newborn. 3rd ed. Philadelphia: Saunders; 1995:373.

4. Curless RG. Subdural empyema in infant meningitis: diagnosis, therapy, and prognosis. Childs Nerv Syst. 1985;1:211.

5. Hobbs C, Childs A-M, Wynne J, et al. Subdural hematoma and effusion in infancy: an epidemiological study. Arch Dis Child. 2005;90:952.

6. Ney JP, Joseph KR, Mitchell MH. Late subdural hygromas from birth trauma. Neurology. 2005;65:517.7. Whitby EH, Griffiths PD, Rutter S, et al. Frequency and natural history of subdural haemorrhages in babies and relation to obstetric factors. Lancet. 2004;363:846.

8. Young S, MacMahon P, Kovar IZ. Subdural intravenous fat collection: an unusual complication of central intravenous feeding in the neonate. J Parenter Enteral Nutr. 1989;13:661.

9. Stine MJ, Harris H. Subdural collection of intravenous fat emulsion in a neonate. Complication of central venous catheterization for total parenteral nutrition. Clin Pediatr. 1985;24:40.

10. Barone MA. Pediatric procedures. In: McMillan JA, Feigin RD, DeAngelis CD, et al., eds. Oski's Pediatrics: Principles and Practice. 4th ed. Philadelphia: Lippincott Williams & Wilkins; 2006:2671.

11. Ferriero D, Buescher ES. Central nervous system. In: Taeusch HW, Christiansen RO, Buescher ES, eds. Pediatric and Neonatal Tests and Procedures. Philadelphia: Saunders; 1996:409.

12. Anand KJ, Johnston CC, Oberlander TF, et al. Analgesia and local anesthesia during invasive procedures in the neonate. Clin Ther. 2005;27:844.

13. O'Brien L, Taddio A, Lyszkiewicz DA, et al. A critical review of the topical local anesthetic amethocaine (Ametop) for pediatric pain. Paediatr Drugs. 2005;7:41.

14. Kaur G, Gupta P, Kumar A. A randomized trial of eutectic mixture of local anesthetics during lumbar puncture in newborns. Arch Pediatr Adolesc Med. 2003;157:1065.

15. Taddio A, Ohlsson A, Einarson TR, et al. A systematic review of lidocaine–prilocaine (EMLA) in the treatment of acute pain in neonates. Pediatrics. 1998;101:e1.

16. Katona F, Balazsm, Berenyi M, et al. Subdural effusion in the first six months of life. Acta Paediatr Acad Sci Hung. 1982;23:219.

17. Gutierrez FA, Raimondi AJ. Acute subdural hematoma in infancy and childhood. Childs Brain. 1975;1:269.

18-Suprapubic Bladder Aspiration

S. Lee Woods

A. Indications (1,2,3,4,5,6,7,8 and 9)

• To obtain urine for culture

Suprapubic bladder aspiration is considered the most reliable method of obtaining urine for culture in infants and children <2 years old. In this age group, the distended bladder is located intraabdominally. Any number of bacteria in urine obtained by this method is considered significant and likely to be indicative of urinary tract infection. Contamination with skin flora can occur, with a false-positive rate of 1% reported (10), but should be avoidable with careful skin preparation. Although bladder catheterization has a higher success rate, it also has a much higher false-positive rate than suprapubic aspiration (11,12). Reported success rates for suprapubic aspiration vary widely, from 25% to 100% (13). With careful attention to performing the procedure when the infant has a full bladder, success is generally 89% to 95%, even in very low-birthweight infants (3,4,7,8 and 9). The use of portable ultrasound (13,14,15,16,17 and 18) or transillumination (19) to determine bladder size can greatly increase the chance of success.

B. Contraindications (1,2,4,6,20)

• Empty bladder as a result of recent void or dehydration

A full bladder is essential for success of the procedure and avoidance of complications.

- Skin infection over the puncture site
- Distention or enlargement of abdominal viscera (e.g., dilated loops of bowel, massive hepatomegaly)
- Genitourinary anomaly or enlargement of pelvic structures (e.g., ovarian cyst, distention of vagina or uterus)
- Uncorrected thrombocytopenia or bleeding diathesis

C. Equipment

All equipment must be sterile, except transillumination light or ultrasound equipment.

- Gloves
- Gauze sponges and cup with iodophor antiseptic solution or
- Prepared antiseptic-impregnated swabs
- 3-mL syringe
- 21- or 22-gauge x $1\hat{A}^{1/2}$ -in needle
- Transillumination light or portable ultrasound (optional)

D. Precautions

- Use strict aseptic technique.
- Delay the procedure if the infant has urinated in the last hour.

If the infant is systemically ill, do not delay antibiotic therapy to wait for further urine production.

• Correct bleeding diathesis before the procedure. Consider catheterization as an alternative.

- Be certain of landmarks. Do not insert the needle over the pubic bone or off the midline.
- Aspirate urine using only gentle suction. The use of too much suction can draw the bladder mucosa to the needle, obstructing the collection of urine and increasing the risk of injury to the bladder.

E. Technique (2,3,4,5 and 6,9,20)

- Have an assistant restrain the infant in the supine, frog-leg position.
- To avoid reflex urination, ask assistant to:
 - Place the tip of a finger in the anus and apply pressure anteriorly in a female infant, or
 - Pinch the base of the penis gently in a male infant.
- Determine the presence of urine in the bladder.
 - Verify that the diaper has been dry for at least 1 hour.
 - Palpate or percuss the bladder.
 - Optionally, use transillumination light (19), or portable ultrasound guidance (13,14,15,16,17 and 18).
- Locate landmarks. Palpate the top of the pubic bone. The site for needle insertion is 1 to 2 cm above the symphysis pubis in the midline (Fig. 18.1).
- Wash hands thoroughly and put on gloves.
- Clean the suprapubic area (including the area over pubic bone) three times with antiseptic solution. Blot dry with sterile gauze.



FIG. 18.1. The bladder in the neonate, with immediate anatomical relations. An asterisk indicates approximate site for needle insertion.

Generally, anesthesia is not required, but local injection of lidocaine at this time or application of topical anesthetic cream prior to cleaning the area can be used for local anesthesia at the puncture sites (21).

• Palpate the symphysis pubis, and insert the needle (with syringe attached) 1 to 2 cm above the pubic symphysis in the midline (Fig. 18.2).



FIG. 18.2. A: Insertion of needle 1 to 2 cm above symphysis publis. B: Midline sagittal section to emphasize the intra-abdominal position of the full bladder in the neonate and its posterior anatomic relations.

- Maintain the needle perpendicular to table or directed slightly caudad.
- Advance the needle 2 to 3 cm. A slight decrease in resistance may be felt when the bladder is penetrated.
- Aspirate gently as the needle is slowly advanced until urine enters the syringe. Do not advance the needle more than 1 in.
 - Withdraw the needle if no urine is obtained.
 - Do not probe with the needle or attempt to redirect it to obtain urine.
 - Wait at least 1 hour before attempting to repeat the procedure.
- Withdraw the needle after urine is obtained. Apply gentle pressure over the puncture site with sterile gauze to stop any bleeding.
- Remove the needle and place a sterile cap on the syringe or transfer urine to a sterile container to send for culture.

F. Complications

Minor transient hematuria is the most commonly reported complication, occurring in 0.6% to 10% of cases (3,7,8 and 9). Serious complications are very rare, occurring in 0.2% of cases or less (4,6).

- Bleeding
 - Transient macroscopic hematuria (blood-tinged urine) (3,7,8 and 9)
 - Gross hematuria (7,8 and 9,22,23 and 24)
 - Abdominal wall hematoma (25)
 - Bladder wall hematoma (7,26)
 - Pelvic hematoma (27)
- Infection
 - Abdominal wall abscess (28,29)
 - Sepsis (30,31)
 - Osteomyelitis of pubic bone (32,33)
- Perforation
 - Bowel (28,31,34,35)

• Pelvic organ (34)

References

1. Long SS, Klein JO. Bacterial infections of the urinary tract. In: Remington JS, Klein JO, Wilson CB, Baker CJ, eds. Infectious Diseases of the Fetus and Newborn. 6th ed. Philadelphia: Saunders; 2006:335. 2. Shilkofski N. Procedures. In: Robertson J, Shilkofski N, eds. The Harriet Lane Handbook. 17th ed. St. Louis, MO: Mosby; 2005:73.

3. Buescher ES. Immunology and infectious diseases procedures and tests. In: Taeusch HW, Christiansen RO, Buescher ES, eds. Pediatric and Neonatal Tests and Procedures. Philadelphia: Saunders; 1996:625.

4. Barkemeyer BM. Suprapubic aspiration of urine in very low birth weight infants. Pediatrics. 1993;92:457.

5. Stevens DC, Schreiner RL, Gresham EL. Suprapubic bladder aspiration in the neonate. Perinatol Neonatol. 1981;5:47.

6. Abbott GD, Shannon FT. How to aspirate urine suprapubically in infants and children. Clin Pediatr. 1970;9:277.

7. Saccharow L, Pryles CV. Further experience with the use of percutaneous suprapubic aspiration of the urinary bladder. Pediatrics. 1969;43:1018.

8. Shannon FT, Sepp E, Rose GR. The diagnosis of bacteriuria by bladder puncture in infancy and childhood. Aust Paediatr J. 1969;5:97.

9. Nelson JD, Peters PC. Suprapubic aspiration of urine in premature and term infants. Pediatrics. 1965;36:132.

10. Pierog S, Parthasarathy S, Rojanaphruk S, et al. Incidence of bacteriuria in high-risk neonate. NY State J Med. 1975;75:2152.

11. Austin BJ, Bollard C, Gunn TR. Is urethral catheterization a successful alternative to suprapubic aspiration in neonates? J Paediatr Child Health. 1999;35:34.

12. Tobiansky R, Evans N. A randomized controlled trial of two methods for collection of sterile urine in neonates. J Paediatr Child Health. 1998;43:460.

13. Gochman RF, Karasic RB, Heller MB. Use of the portable ultrasound to assist urine collection by suprapubic aspiration. Ann Emerg Med. 1991;20:6.

14. Chu RW, Wong YC, Luk SH, et al. Comparing suprapubic urine aspiration under real time ultrasound guidance with conventional blind aspiration. Acta Paediatr. 2002;91:512.

15. Munir V, Barnett P, South M. Does the use of volumetric bladder ultrasound improve the success rate of suprapubic aspiration of urine? Pediatr Emerg Care. 2002;18:346.

16. Ozkan B, Kaya O, Akdag R, et al. Suprapubic bladder aspiration with or without ultrasound guidance. Clin Pediatr (Phila.). 2000;39:625.

17. Garcia-Neito VG, Navarro JF, Sanchez-Almeida ES, et al. Standards for ultrasound guidance of suprapubic bladder aspiration. Pediatr Nephrol. 1997;11:607.

18. Kiernan SC, Pinckert TL, Keszler M. Ultrasound guidance of suprapubic bladder aspiration in neonates. J Pediatr. 1993;123:789.

19. Buck JR, Weintraub WH, Coran AG, et al. Fiberoptic transillumination: a new tool for the pediatric surgeon. J Pediatr Surg. 1977;12:451.

20. Barone MA. Pediatric procedures. In: McMillan JA, Feigin RD, DeAngelis CD, et al., eds. Oski's Pediatrics: Principles and Practice. 4th ed. Philadelphia: Lippincott Williams & Wilkins; 2006:2671.

21. Anand KJ, Johnston CC, Oberlander TF, et al. Analgesia and local anesthesia during invasive procedures in the neonate. Clin Ther. 2005;27:844.

22. Carlson KP, Pullon DHH. Bladder hemorrhage following transcutaneous bladder aspiration. Pediatrics. 1977;60:765.

23. Lanier B, Daeschner CW. Serious complication of suprapubic aspiration of the urinary bladder. J Pediatr. 1971;79:711.

24. Rockoff AS. Hemorrhage after suprapubic bladder aspiration. J Pediatr. 1976;89:327.

25. Kunz HH, Sieberth HG, Freiberg J, et al. Zur Bedeutung der Blasenpunktion fur den sicheren Nachweis einer Bakteriurie. Dtsch Med Wochenschr. 1975;100:2252.

26. Morell RE, Duritz G, Oltorf C. Suprapubic aspiration associated with hematoma. Pediatrics. 1982;69:455.

27. Mandell J, Stevens P. Supravesical hematoma following suprapubic urine aspiration. J Urol. 1978;119:286.

28. Polnay, L, Fraser AM, Lewis JM. Complication of suprapubic bladder aspiration. Arch Dis Child. 1975;50:80.

29. Uhari M, Remes M. Suprapubic abscessâ€"a complication of suprapubic bladder aspiration. Arch Dis Child. 1977;52:985.

30. Mustonen A, Uhari M. Is there bacteremia after suprapubic aspiration in children with urinary tract infection? J Urol. 1978;119:822.

31. Pass RF, Waldo FB. Anaerobic bacteremia following suprapubic bladder aspiration. J Pediatr. 1979;94:748.

32. Wald ER. Risk factors for osteomyelitis. Am J Med. 1985;78:206.

33. Kalager T, Digranes A. Unusual complication after suprapubic bladder puncture. Br Med J. 1979;1:91.

34. Weathers WT, Wenzel JE. Suprapubic aspiration: perforation of a viscus other than the bladder. Am J Dis Child. 1969;117:590.

35. Schreiner RL, Skafish P. Complications of suprapubic bladder aspiration. Am J Dis Child. 1978;132:98.

19-Bladder Catheterization

S Lee Fragneto

A. Indications (1,2,3,4,5,6 and 7)

• To obtain urine for culture, particularly when suprapubic collection is contraindicated and when clean-catch specimen is unsatisfactory

While suprapubic bladder aspiration is considered the most reliable method of obtaining urine for culture in infants and young children (see Chapter 18), bladder catheterization is an acceptable alternative method (5). Bladder catheterization has a higher success rate than suprapubic aspiration, especially if the practitioner is inexperienced in bladder aspiration. However, urine samples collected by catheterization have a higher false-positive rate than suprapubic aspiration (3,5,8,9 and 10), and catheterization can introduce bacteria colonizing the distal urethra into the bladder, causing a urinary tract infection (see F, $\hat{a}\in \infty$ Complications $\hat{a}\in$). The diagnosis of urinary tract infection cannot be made reliably by culturing urine collected in a bag (5,11).

- To monitor precisely the urinary output of a critically ill patient
- To quantify bladder residual
- To relieve urinary retention (e.g., in neurogenic bladder) (12,13 and 14)
- To instill contrast agent to perform cystourethrography (15)

B. Contraindications

In the presence of uncorrected bleeding diathesis, potential risks and benefits must be considered. C. Equipment

All equipment must be sterile. Commercial prepackaged urinary drainage kits, with or without collection burettes for closed drainage, are available.

- Gloves
- Gauze sponges and cup with iodophor antiseptic solution (not containing alcohol), or

- Prepared antiseptic-impregnated swabs
- Towels for draping
- Surgical lubricant
- Cotton-tipped applicators
- Urinary catheter

Silicone urinary drainage catheters are available in 3.5, 5.0, 6.5, and 8.0 French (Fr) sizes. A 5-Fr infant feeding tube or a 3.5- or 5.0-Fr umbilical catheter may be substituted for a urinary catheter.

• Sterile container for specimen collection or collection burette for continuous closed drainage

D. Precautions

- Use strict aseptic technique.
- Use adequate lighting.
- Try to time the procedure for when the infant has not recently voided (1 to 2 hours after the last wet diaper). Portable ultrasound can be helpful in determining when there is sufficient urine present in the bladder, reducing the chance of an unsuccessful attempt (16,17).
- Avoid vigorous irrigation of the perineum in preparation for catheterization. This may increase the risk of introducing bacteria into the urinary tract.
- Avoid separating the labia minora too widely, to prevent tearing of the fourchette.
- Use the smallest-diameter catheter to avoid traumatic complications. A 3.5-Fr catheter is recommended for infants weighing less than 1,000 g and a 5-Fr catheter is recommended for larger infants.
- If the catheter does not pass easily, do not use force. Suspect obstruction and abandon the procedure.
- To avoid coiling and knotting, insert the catheter only as far as necessary to obtain urine.
- If urine is not obtained in a female infant, recheck the location of the catheter by visual inspection or by radiographic examination. It may have passed through the introitus into the vagina.
- Remove the catheter as soon as possible to avoid infectious complications.
- If the catheter cannot be removed easily, do not use force. Consult urology, as it may be knotted.

E. Technique

Male Infant (2,4,18,19)

- Set up equipment and squeeze a small amount of lubricant onto a sterile field.
- Restrain the infant supine in the frog-leg position.
- Wash hands thoroughly and put on gloves.
- Stabilize the shaft of the penis with the nondominant hand. This hand is now considered contaminated.
- If the infant is uncircumcised, gently retract the foreskin just enough to expose the meatus. Do not attempt to lyse adhesions. The young male infant has physiologic phimosis, and the foreskin cannot be fully retracted (19,20). If the foreskin is tightly adherent, attempt to line up the preputial ring and the meatus.
- Apply gentle pressure at the base of the penis to avoid reflex urination.
- Using the free hand for the rest of the procedure, clean the glans three times with antiseptic solution. Begin at the meatus and work outward and down the shaft of the penis. Blot dry with sterile gauze.
- Drape sterile towels across the lower abdomen and across the infant's legs.
- Place the wide end of the catheter or feeding tube into the specimen container.
- Lubricate the tip of the catheter copiously.
- Move the specimen container and catheter onto the sterile drape between the infant's legs.
- Gently insert the catheter through the meatus just until urine is seen in the tube (Fig. 19.1).

- During insertion, apply gentle upward traction on the penile shaft to prevent kinking of the urethra (Fig. 19.1).
- If the meatus cannot be visualized, insert the catheter through the preputial ring in a slightly inferior direction. If there is any question about catheter position, abandon the procedure.
- If resistance is met at the external sphincter, hold the catheter in place, applying minimal pressure. Generally, spasm will relax after a brief period, allowing easy passage of catheter. If not, suspect obstruction and abandon the procedure.
- Do not move the catheter in and out. This will increase the risk of urethral trauma.
- Do not insert extra tubing length in an attempt to stabilize a catheter to be left indwelling. This will increase the risk of trauma and knotting.
- Collect specimen for culture.
- If the catheter is to remain indwelling, connect the catheter immediately to a closed sterile system for urine collection. Tape the tube securely to the inner thigh.
- If the catheter is to be removed, gently withdraw it when urine flow ceases.



FIG. 19.1. Anatomic drawing demonstrating bladder catheterization in the male.

Female Infant (7,19,21)

- Follow Steps 1 through 3 of technique for male infant.
- Retract the labia minora.
 - Use sterile gauze sponges with nondominant hand, or
 - Have an assistant retract the labia with two cotton-tipped applicators (Fig. 19.2).
- Using the free hand for the rest of the procedure, cleanse the area between the labia minora three times with antiseptic solution.
 - Swab in an anterior-to-posterior direction to avoid drawing fecal material into the field.
 - Blot dry with sterile gauze.
- Follow Steps 8 through 11 of the technique for the male infant.
- Visualize the meatus (Fig. 19.2).
 - The most prominent structure is the vaginal introitus. The urethral meatus lies immediately anterior (between the clitoris and the introitus).
 - The meatus may be obscured by the introital fold. Gently push the fold down with a cotton-tipped applicator.

- If the meatus is not visible, the infant may have female hypospadias (the meatus is on the roof of the vagina, just inside the introitus) (19). The urethra must then be catheterized blindly, which may require a curved-tip catheter or urologic assistance.
- Gently insert the catheter only until urine appears in the tube. Do not insert extra tubing.
- Follow Steps 13 through 15 of technique for the male infant.

Female Infant in Prone Position (22)

This technique is useful in an infant who cannot be placed supine (e.g., one with a large meningomyelocele).

- Position the infant prone on folded blankets so the head and trunk are elevated about 3 in above the knees and lower legs. The hips should be flexed with knees abducted (Fig. 19.3A).
- Place a gauze pad over the anus and secure with tape across the buttocks, to avoid contamination of the perineum from reflex bowel evacuation (Fig. 19.3B).
- Place sterile drapes as shown in Fig. 19.3C. Follow the procedure for female catheterization above.



FIG. 19.2. External genitalia in the female. Retraction of labia majora and minora with cotton-tipped applicators. An arrow indicates urethral meatus.

F. Complications

- Infection (23,24,25,26 and 27)
 - Urethritis
 - o Epididymitis
 - Cystitis
 - o Pyelonephritis
 - o Sepsis

The most common complication of bladder catheterization is the introduction of bacteria into the urinary tract and potentially into the bloodstream. Catheterization is the leading cause of nosocomial urinary tract infection and gram-negative sepsis in adult patients (24). The risk of bacteriuria from straight ($\hat{a}\in \alpha$ in-and-out $\hat{a}\in$) catheterization is 1% to 5% in this population (23,24 and 25). The risk of infection is related directly to the duration of catheterization. In infants and children, approximately 50% to 75% of hospital-acquired urinary tract infections occur in catheterized patients, the highest rate being in neonates (26,27). Urinary tract

infection developed in 10.8% of catheterized pediatric patients (26), and secondary bacteremia in 2.9% (27). Risk of infection is decreased by adhering to strict aseptic technique during catheter placement, maintaining a closed sterile collection system, and removing the catheter as soon as possible.



FIG. 19.3. A: Position of infant for prone catheterization. B: Placement of gauze pad over anus. C: Placement of drapes. (Adapted from

Campbell J. Catheterizing prone female infants: how can you see what you're doing? Am J Matern Child Nurs. 1979;4:376, based on drawing by N. L. Gahan, with permission.)

- Trauma
 - o Hematuria
 - Urethral erosion or tear (28)
 - Urethral false passage (28,29)
 - Perforation of the urethra or bladder (Fig. 19.4) (28,30,31)
 - Tear of the fourchette (28)
 - Meatal stenosis (20)
 - Urethral stricture (32)
 - Urinary retention secondary to urethral edema (28)

The risk of trauma is reduced by using the smallest-diameter catheter with ample lubrication, advancing the catheter only as far as necessary to obtain urine, and never forcing a catheter through an obstruction. Erosion and perforation are associated with long-indwelling catheters. This risk is reduced by removing the catheter as soon as possible.

- Mechanical
 - Catheter malposition (19,28)
 - Catheter knot (33,34,35,36 and 37)

The risk of knotting is reduced by using the minimal length of catheter insertion. Standard insertion lengths of 6 cm for male and 5 cm for female term newborns have been suggested (37). Shorter lengths would be appropriate for preterm infants. A more general standard is to insert the catheter only as far as needed to obtain urine. Using a feeding tube as a urinary

catheter may also increase the risk of knotting, because these tubes are softer and more likely to coil.



FIG. 19.4. A: Cystogram shows dilated posterior urethra (arrows) secondary to posterior urethral valves. B: Subsequent film shows perforation of the bladder, with free contrast material in the peritoneal cavity.

References

1. Long SS, Klein JO. Bacterial infections of the urinary tract. In: Remington JS, Klein JO, Wilson CB, Baker CJ, eds. Infectious Diseases of the Fetus and Newborn. 6th ed. Philadelphia: Saunders; 2006:335. 2. Shilkofski N. Procedures. In: Robertson J, Shilkofski N, eds. The Harriet Lane Handbook. 17th ed. St. Louis, MO: Mosby; 2005:73.

3. Ma JF, Diariki Shortliffe LM. Urinary tract infection in children: etiology and epidemiology. Urol Clin NA. 2004;31:517.

4. Carter HB. Basic instrumentation and cystoscopy. In: Walsh PC, Retik AB, Vaughan ED, et al., eds. Campbell's Urology. 8th ed. Philadelphia: Saunders; 2002:11.

5. Committee on Quality Improvement, Subcommittee on Urinary Tract Infection. Practice parameter: the diagnosis, treatment, and evaluation of the initial urinary tract infection in febrile infants and young children. Pediatrics. 1999;103:843.

6. Buescher ES. Immunology and infectious diseases procedures and tests. In: Taeusch HW, Christiansen RO, Buescher ES, eds. Pediatric and Neonatal Tests and Procedures. Philadelphia: Saunders; 1996:625.7. Redman JF, Bissada NK. Direct bladder catheterization in infant females and young girls. Clin Pediatr. 1976;15:1060.

8. Austin BJ, Bollard C, Gunn TR. Is urethral catheterization a successful alternative to suprapubic aspiration in neonates? J Paediatr Child Health. 1999;35:34.

9. Tobiansky R, Evans N. A randomized controlled trial of two methods for collection of sterile urine in neonates. J Paediatr Child Health. 1998;43:460.

10. Pollack CV, Pollack ES, Andrew ME. Suprapubic bladder aspiration versus urethral catheterization in ill infants: success, efficiency and complication rates. Ann Emerg Med. 1994;23:225.

11. Al-Orifi F, McGillivray D, Tange S, et al. Urine culture from bag specimens in young children: are the risks too high? J Pediatr. 2000;137:221.

12. Ewalt DH, Bauer SB. Pediatric neurourology. Urol Clin North Am. 1996;23:501.

13. Baskin LS, Kogan BA, Benard F. Treatment of infants with neurogenic bladder dysfunction using anticholinergic drugs and intermittent catheterisation. Br J Urol. 1990;66:532.

14. Joseph DB, Bauer SB, Colodny AH, et al. Clean intermittent catheterization of infants with neurogenic bladder. Pediatrics. 1989;84:78.

15. Shalaby-Rana E, Lowe LH, Blask AN, et al. Imaging in pediatric urology. Pediatr Clin North Am. 1997;44:1065.

16. Milling TJ, Van Amerongen R, Melville L, et al. Use of ultrasonography to identify infants in whom urinary catheterization will be unsuccessful because of insufficient urine volume: validation of the urinary bladder index. Ann Emerg Med. 2005;45:510.

17. Chen L, Hsiao AL, Moore L, et al. Utility of bedside bladder ultrasound before urethral catheterization in young children. Pediatrics. 2005;115:108.

 Barone MA. Pediatric procedures. In: McMillan JA, Feigin RD, DeAngelis CD, et al., eds. Oski's Pediatrics: Principles and Practice. 4th ed. Philadelphia: Lippincott Williams & Wilkins; 2006:2671.
Ben-Ami T, Lebowitz RL. Pediatric uroradiology. In: Retik AB, Cukier J, eds. Pediatric Urology.

Baltimore: Williams & Wilkins; 1986:32.

20. Brown MR, Cartwright PC, Snow BW. Common office problems in pediatric urology and gynecology. Pediatr Clin North Am. 1997;44:1091.

21. Redman JF. Techniques of genital examination and bladder catheterization in female children. Urol Clin North Am. 1990;17:1.

22. Campbell J. Catheterizing prone female infants: how can you see what you're doing? Am J Matern Child Nurs. 1979;4:376.

23. Nadler BB, Bushman W, Wyker AW. Standard diagnostic considerations. In: Gillenwater JY, Grayhack JT, Howard SS, et al., eds. Adult and Pediatric Urology. Philadelphia: Lippincott, Williams & Wilkins; 2002:47.

24. Sedor J, Mulholland SG. Hospital-acquired urinary tract infections associated with the indwelling catheter. Urol Clin North Am. 1999;26:821.

25. Warren JW. Clinical presentation and epidemiology of urinary tract infections. In: Mobley HLT, Warren JW, eds. Urinary Tract Infections: Molecular Pathogenesis and Clinical Management. Washington, DC: ASM Press; 1996:3.

26. Lohr JA, Downs SM, Dudley S, et al. Hospital-acquired urinary tract infections in the pediatric patient: a prospective study. Pediatr Infect Dis J. 1994;13:8.

27. Dele Davies H, Ford Jones EL, Sheng RY, et al. Nosocomial urinary tract infections at a pediatric hospital. Pediatr Infect Dis J. 1992;11:349.

28. McAlister WH, Cacciarelli A, Shackelford GD. Complications associated with cystography in children. Radiology. 1974;111:167.

29. Koleilat N, Sidi AA, Gonzalez R. Urethral false passage as a complication of intermittent catheterization. J Urol. 1989;142:1216.

30. Basha M, Subhani M, Mersal A, et al. Urinary bladder perforation in a premature infant with Down syndrome. Pediatr Nephrol. 2003;18:1189.

31. Salama H, Al Ju Fairi M, Rejjal A, et al. Urinary bladder perforation in a very low birth weight infant. J Perinat Med. 2002;30:440.

32. Edwards LE, Lock R, Powell C, et al. Post-catheterisation urethral strictures. A clinical and experimental study. Br J Urol. 1983;55:53.

33. Anatol T, Nunez J. Intravesical tube knot in a neonate. J Trop Pediatr. 2005;51:314.

34. Lodha A, Ly L, Brindle M, et al. Intraurethral knot in a very-low-birth-weight infant: radiological recognition, surgical management and prevention. Pediatr Radiol. 2005;35:713.

35. Anbu AT, Palmer K. Urethral catheter knotting in preterm neonates. Indian Pediatr. 2004;41:631.36. Mayer E, Ankem MK, Hartanto VH, et al. Management of urethral catheter knot in a neonate. Can J Urol. 2002;9:1649.

37. Carlson D, Mowery BD. Standards to prevent complications of urinary catheterization in children: should and should-knots. J Soc Pediatr Nurs. 1997;2:37.

20-Tympanocentesis

Hosai Hesham Gregory J. Milmoe

A. Indications

Diagnostic tympanocentesis is indicated in neonatal acute otitis media (AOM) in order to target antibiotic therapy. Myringotomy may be used for both diagnostic and drainage purposes. The specific indications include:

- AOM not responding to antibiotics after 72 hours
- AOM in severely immunocompromised infant
- AOM in infant already on antibiotics
- AOM with suppurative complications, e.g., mastoiditis, facial paralysis, sepsis
- To confirm the diagnosis when the clinical exam is not clear
- To relieve severe otalgia

B. Contraindications

- Difficulty in confirming ossicular landmarks. One must be able to identify the malleus and the annulus of the tympanic membrane (TM) (Fig. 20.1).
- Suggestion of abnormal anatomy. This may be more likely in patients with congenital malformation syndromes.
- Suggestion of alternate pathology, e.g., cholesteatoma or neoplasm

C. Equipment

- Surgical gloves
- Otoscope with open operating head and good light
- Largest speculum that will fit the canal (2, 3, or 4 mm)
- 18-gauge 3-in spinal needle with 1-mL or 3-mL syringe
- Blunt ear curette
- 70% isopropyl alcohol in 3-mL syringe for cleaning and antisepsis of ear canal
- Suction setup with No. 5 Frazier ear suction
- Culturettes with transport media

D. Precautions

- Patient safety and comfort require proper restraint, adequate light, and appropriate instruments.
- The kindest way is to be quick, and this means having the child still.
- Conscious sedation is feasible only if the child is stable and has no issues with airway obstruction. It is not needed past the point of puncturing the TM, so most often no medicine is used.
- Good visualization is paramount. Sufficient cleaning must be done so that the malleus and the anterior aspect of the annulus are clearly seen.
- Avoid the posterior aspect of the tympanic membrane. This is where the round window, stapes, and incus are present.

E. Technique

- Restrain child fully with hospital linen wrapped around the torso and arms.
- Position child with the head turned so that the involved ear is up. The assistant must keep the head still.
- Rinse ear canal with alcohol solution from a 3-mL syringe. This will provide antisepsis and initiate cleaning.
- Let fluid run out or use suction.
- Use otoscope to visualize canal and remove debris with curette or suction.
- Align speculum to get best view of TM landmarks. Pulling superiorly and laterally on the pinna will help immensely (Fig. 20.2).
- Attach spinal needle to syringe after bending it 45 to 60 degrees at the hub. This keeps the syringe out of your line of sight.
- Hold needle at the hub and introduce it through the otoscope. Puncture the drum anterior to the malleus at or below the umbo level (Fig. 20.3).
- Hold needle securely and have assistant draw back on the syringe to obtain the sample.
- Place sample in appropriate transport medium.
- If more drainage is required, a myringotomy blade can be used to widen the opening. This will still close in 48 to 72 hours.



FIG. 20.1. Normal newborn eardrum. View through speculum.

F. Complications

- Most common is bleeding from canal wall. This will stop but is preferably avoided.
- TM perforation that persists. Initially this is actually helpful for drainage and ventilation of the middle ear space.
- Disruption of the ossicles from malpositioned needle (see D, Precautions)
- Major bleeding from dehiscent jugular bulb or carotid artery rare (see D, Precautions, and B, Contraindications)



FIG. 20.2. Tympanic membrane in the adult (A) and infant (B). The portion of the tympanic membrane that may be visualized through the speculum at one time is within the dotted line.



FIG. 20.3. Tympanocentesis. Aspirating the middle ear using a 3-mL syringe. Needle is penetrating eardrum inferiorly.

References

1. Burton DM, Seid AB, Kearns D, et al. Neonatalotitis media. Arch Otolaryngol Head Neck Surg. 1993;119:672–675.

2. Arriaga MA, Bluestone CD. The role of tympanocentesis in the management of infants with sepsis. Laryngoscope. 1989;99: 1048–1051.

3. Nomura Y, Mimata H, Yamasaki M, et al. Effect of myringotomy on prognosis in pediatric acute otitis media. Int J Pediatr Otorhinolaryngol. 2005;69:61–64.

4. Guarisco JL, Grundfast KM. A simple device for tympanocentesis in infants and children. Laryngoscope. 1988;98: 244.

5. Bluestone CD, Klein JO. Otologic surgical procedures. In: Bluestone CD, Stool SE, Kenna M, eds. Pediatric Otolaryngology. Philadelphia: Saunders; 1996:28–35.

6. Hasebe S, et al. Proximity of carotid canal wall to tympanic membrane: a human temporal bone study. Laryngoscope. 2003;113: 802–807.

21-Tibial Bone Marrow Biopsy

Martha C. Sola-Visner Lisa M. Rimsza Robert D. Christensen

A. Purpose

- To obtain a bone marrow clot sample for histologic evaluation of the following¹
 - Bone marrow cellularity
 - Relative abundance of myeloid, erythroid, lymphoid, and megakaryocytic lineages, using specific immunohistochemical stains on multiple cuts if necessary
 - \circ $\,$ Maturation and morphology of cells of all lineages $\,$
 - Presence of infiltrative nonmalignant diseases
 - Presence of infiltrative malignant diseases (hematologic and nonhematologic)
 - Presence of granulomas or infectious organisms
- To obtain emergent cytogenetic studies

B. Indications

- Evaluation of hematologic disorders (1,2,3,4,5,6)
 - Suspected neonatal aplastic anemia (pancytopenia)
 - Suspected leukemia, when blood studies are insufficient to confirm the diagnosis
 - Neutropenia of unclear etiology, which is severe (absolute neutrophil count <500/mL) and persists for more than 1 week
 - Severe neutropenia (<500/mL), which is unresponsive to 3 days of treatment with recombinant granulocyte colony-stimulating factor
 - Neutropenia of unclear etiology, which is moderately severe (500 to 999/mL) and persists for more than 2 weeks
 - Thrombocytopenia of unclear etiology, which is severe (platelets <50,000/mL) and persists for more than 1 week
- Evaluation of suspected metabolic storage disease (e.g., Niemann-Pick disease) (2)
- Evaluation of suspected hemophagocytic syndrome or familial hemophagocytic lymphohistiocytosis (7,8)
- Detection of infiltrating tumor cells (9,10,11,12) or of congenital systemic Langerhans' cell histiocytosis (13)
- Certain cultures, e.g., in disseminated tuberculosis or fungal disease (14)
- Cytogenetic studies, for chromosomal analysis (even after transfusion of donor blood) within 3 to 4 hours (15)

C. Contraindications

- Sampling from the sternum is not recommended because of danger of damage to intrathoracic and mediastinal organs (2,16).
- Sampling from the anterior iliac crest is not recommended, particularly in the smallest preterm infants, owing to the proximity to intra-abdominal organs.
- Risks/benefits should be considered carefully in the presence of coagulopathy or when administering anticoagulants or thrombolytics.
- Risks/benefits should be carefully considered in preterm infants with severe osteopenia of prematurity (17).

D. Limitations

• In very small preterm infants, the tibial bone marrow biopsy technique sometimes yields no marrow or a very hemodilute sample, mostly because of the small size of the marrow compartment within the tibia.

E. Equipment Sterile

- Surgical gloves
- Cup with antiseptic solution
- Gauze squares
- Sterile drapes
- 1% lidocaine without epinephrine in 1-mL syringe, with 27-gauge needle
- 19-gauge, Â¹/₂-in Osgood bone marrow needle (Popper and Sons, New Hyde Park, NY, USA) (Fig. 21.1)
- 3-mL syringe without Luer-Lok

Nonsterile

- Cup containing fixative
- 1- to 2-in needle to aid in removal of clot from the syringe



FIG. 21.1. View of the 19-gauge, $\hat{A}^{1/2}$ -in Osgood bone marrow needle. The trocar must be completely inserted in the Osgood needle prior to the procedure.

F. Precautions

- Correct coagulopathy as far as possible prior to procedure.
- Use a total of 0.2 to 0.4 mL of lidocaine. Aspirate before injection to avoid intravascular injection.
- Stabilize the leg in your hand, between your thumb and forefinger. To avoid bone fracture, be sure to apply counterpressure with your palm directly opposite the site of penetration.
- Be aware that less pressure is required to insert the bone marrow needle in neonates (particularly in very low-birthweight infants) than in older children.

- Be careful to enter the bone 1 to 2 cm below the tibial tuberosity, to minimize the risk of injuring the growth plate.
- After the procedure, apply adequate pressure to control bleeding.

G. Special Circumstances

• In cases of suspected osteopetrosis, obtaining an iliac crest bone/bone marrow biopsy is preferable, because it allows quantification of osteoclasts and evaluation of marrow and bony changes consistent with osteopetrosis. In these cases, the tibial bone marrow biopsy technique usually yields only blood or no sample.

H. Technique

- Place the infant in the supine position.
- Use the triangular area at the proximal end of the medial (flat) surface of the tibia, approximately 1 to 2 cm distal to the tibial tuberosity (18).
- Prepare and drape as for a major procedure (see Chapter 4).
- Infiltrate subcutaneous tissue with lidocaine as the needle is slowly advanced. Inject further small volume when the needle reaches the bone, making sure that the tip of the needle is inserted into the bone for subperiosteal injection.
- Remove the needle and wait 2 to 3 minutes.
- Use your nondominant hand to firmly stabilize the leg, providing support with your palm directly opposite the site of marrow puncture. This hand cannot be reintroduced into the sterile field.
- Make sure that the trocar is completely inserted in the Osgood needle.
- Hold the needle between the thumb and forefinger of your dominant hand.
- Introduce the needle at a 90-degree angle, and advance it into the marrow cavity with a slow, twisting motion (Fig. 21.2).
- Continue to advance the needle until it is firmly fixed in bone (does not move when touched) (Fig. 21.3).
- Remove the trocar from the needle and advance the hollow needle an additional 2 to 3 mm into the marrow space (this trephinates marrow spicules into the needle).
- Attach a 3-mL syringe (without a Luer-Lok) firmly to the needle.
- Withdraw the plunger forcefully until a small drop of marrow (<0.05 mL) appears in the syringe hub. Suction should be stopped as soon as the smallest amount of marrow is obtained, because excessive suction will dilute the sample with peripheral blood.
- If no marrow is obtained initially, rotate, advance, or retract the needle and try again.
- Remove the syringe as soon as bone marrow is obtained and withdraw the plunger (with marrow attached) to the bottom of the syringe. Allow the marrow to clot there.
- Remove the needle and apply pressure over the site to achieve hemostasis.

Preparation of the Bone Marrow Clot

- Once the marrow specimen has clotted, dislodge the clot gently with the use of a 1- or 2-in needle and place it into the fixative solution (Fig. 21.4).
- Process the bone marrow clot in a manner identical to a typical bone marrow biopsy, except that decalcification is not required (Fig. 21.5).



FIG. 21.2. The Osgood needle is introduced into the tibial marrow cavity with a slow, twisting motion. Notice that the leg is firmly stabilized in the operator's nondominant hand.



FIG. 21.3. The Osgood needle is firmly fixed in the bone.

I. Complications²

- Subperiosteal bleeding (19)
- Cellulitis or osteomyelitis (20)
- Limb fracture (21)
- Injury to blood vessels (19)
- Bone changes on x-ray film (22,23)
 - Lytic lesions
 - Exostoses
 - Subperiosteal calcification (secondary to hematoma)



FIG. 21.4. A small amount of bone marrow has been obtained in a 3-mL syringe and allowed to clot at the bottom of the syringe. The plunger has been removed, and the clot is now being gently dislodged from the plunger (with the use of a 1- or 2-in needle) and placed into the fixative solution.



FIG. 21.5. Photomicrograph of a bone marrow clot section obtained from a neutropenic neonate. The cellularity is near 100%. Myeloid precursors, scattered erythroid cells, lymphocytes, and several megakaryocytes are clearly identified. Hematoxylin and eosin; original magnification x200.

- J. Advantages of the Tibial Site
 - It is a safe site, particularly in very small preterm infants, because it avoids any proximity to vital organs.
 - The tibia can be easily positioned without disturbing even the sickest infants (usually maintained in the supine position while on mechanical ventilation).
 - It can be adequately stabilized and supported by the nondominant hand of the person performing the procedure.

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References

1. Calhoun DA, Christensen RD, Edstrom CS, et al. Consistent approaches to procedures and practices in neonatal hematology. Clin Perinatol. 2000;27:733.

2. Downing V. Bone marrow examination in children. Pediatr Clin North Am. 1955;2:243.

3. Garcia L, Valcarcel M, Santiago-Borrero PJ. Chemotherapy during pregnancy and its effects on the fetusâ€"neonatal myelosuppression: two case reports. J Perinatol. 1999;19:230.

4. Juul SE, Calhoun DA, Christensen RD. "Idiopathic neutropenia†in very-low birthweight infants. Acta Paediatr. 1998;87:963.

5. Calhoun DA, Kirk JF, Christensen RD. Incidence, significance, and kinetic mechanism responsible for leukemoid reactions in patients in the neonatal intensive care unit: a prospective evaluation. J Pediatr. 1996;129:403.

6. Mizutani K, Azuma E, Komada Y, et al. An infantile case of cytomegalovirus induced idiopathic thrombocytopenic purpura with predominant proliferation of CD10 positive lymphoblasts in bone marrow. Acta Paediatr Jpn. 1995;37:71.

7. Aygun C, Tekinalp G, Gurgey A. Infection-associated hemophagocytic syndrome due to Pseudomonas aeruginosa in preterm infants. J Pediatr Hematol Oncol. 2003;25:665.

8. Rugolotto S, Marradi PL, Balter R, et al. Familial haemophagocytic lymphohistiocytosis: survival of a premature twin with immuno-chemotherapy and bone marrow transplantation from an HLA-identical unrelated donor. Acta Paediatr. 2005;94:971.

9. Karcioglu ZA, Al-Mesfer SA, Abboud E, et al. Workup for metastatic retinoblastoma. A review of 261 patients. Ophthalmology. 1997;104:307.

10. Moscinski LC, Pendergrass TW, Weiss A, et al. Recommendations for the use of routine bone marrow aspirations and lumbar punctures in the follow up of patients with retinoblastoma. J Pediatr Hematol Oncol. 1996;18:130.

11. Osmanagaoglu K, Lippens M, Benoit Y, et al. A comparison of iodine-123 meta-iodobenzylguanidine scintigraphy and single bone marrow aspiration biopsy in the diagnosis and follow-up of 26 children with neuroblastoma. Eur J Nucl Med. 1993;20:1154.

12. Penchansky L. Bone marrow biopsy in the metastatic work-up of solid tumors in children. Cancer. 1984;54:1447.

13. Stiakaki E, Giannakopoulou C, Kouvidi E. Congenital systemic Langerhans cell histiocytosis (report of two cases). Haematologia (Budap). 1997;28:215.

14. Machin GA, Honore LH, Fanning EA, et al. Perinatally acquired neonatal tuberculosis: report of two cases. Pediatr Pathol. 1992;12:707.

15. Page BM, Coulter JB. Bone marrow aspiration for chromosome analysis in newborn. Br Med J. 1978;1:1455.

16. Bakir F. Fatal sternal puncture. Dis Chest. 1963;44:435.

17. Dabezies EJ, Warren PD. Fractures in very low birth weight infants with rickets. Clin Orthop Relat Res. 1997;335:233.

18. Sola MC, Rimsza LM, Christensen RD. A bone marrow biopsy technique suitable for use in neonates. Br J Haematol. 1999;107:458.

19. McNutt DR, Fudenberg HH. Bone-marrow biopsy and osteoporosis. N Engl J Med. 1972;1:46.

20. Shah M, Watanakunakorn C. Staphylococcus aureus sternal osteomyelitis complicating bone marrow aspiration. South Med J. 1978;71:348.

21. Miller D. Normal values and examination of the blood: perinatal period, infancy, childhood, and adolescence. In: Miller D, Pearson H, Bachner R, et al., eds. Smith's Blood Diseases of Infancy and Childhood. St. Louis, MO: Mosby; 1978:20–21.

22. Gilsanz V, Grunebaum M. Radiographic appearance of iliac marrow biopsy sites. AJR. 1977;128:597. 23. Murphy WA. Exostosis after iliac bone marrow biopsy. AJR. 1977;129:1114.

22-Punch Skin Biopsy

Pamela Jakubowicz

A. Definition

• A small, full-thickness biopsy utilizing a cylindrical instrument

B. Indications

- Diagnosis of skin lesions (1,2,3,4,5,6,7 and 8)
- Electron and light microscopic identification of certain hereditary and metabolic disorders (9,10,11,12,13,14 and 15)
- Genetic, enzymatic, or morphologic studies on established fibroblast strains (16)
- Treatment of small skin lesions

C. Types of Skin Biopsy (6,10,17)

- Punch skin biopsy is appropriate when full thickness is necessary.
 - Allows for pathologic evaluation and rapid diagnosis of certain conditions
- Incisional biopsies are used predominantly for disorders of deep subcutaneous fat or fascia (e.g., erythema nodosum).
- Excision of larger lesions by a trained dermatologist or surgeon is preferable when planning to remove an entire large lesion.

D. ContraindicationsBleeding disorder when risk outweighs benefitsE. EquipmentSterile

- Towel or tray to form sterile area
- 70% alcohol or other suitable antiseptic agent
- 4- x 4-in gauze squares
- Lidocaine HCl 1% with epinephrine in 1-mL tuberculin syringe with 27- or 30-gauge needle (use epinephrine-free local anesthesia on extremeties to avoid distorting skin vascularity in patients with urticaria pigmentosa)
- Blunt tissue forceps
- Fine, curved scissors or no. 15 scalpel blade
- Sharp 3- or 4-mm punch¹ (Fig. 22.1)

Skin biopsy has been performed on the fetus (11,18) and may be done postmortem on stillborn or recently deceased infants to produce fibroblast cultures for karyotype (see Chapter 24). Under the latter circumstances, punch or excisional biopsy from the freshest-appearing, least-macerated skin area(s) is appropriate.

- 5-0 or 6-0 nylon suture with small curved needle on needle holder, Dermabond (Ethicon, Somerville, NJ, USA) or Steri-Strip (3M Health Care, St. Paul, MN, USA)
- Adhesive bandage with petrolatum jelly
- Appropriate transport medium affixed with patient's information (Table 22.1) (In addition, a razor may be required for hairy areas)

F. Precautions

- Avoid sites, if possible, where a small scar would potentially be cosmetically disfiguring.
 - Tip, bridge, and columella of nose
 - o Eyelids
 - Lip margins
 - \circ Nipples
 - \circ Fingers or toes
 - Areas overlying joints
 - Lower leg below the knee
- Avoid very small punch (2 mm or less), because this limits ability to interpret pathologic findings.
- Avoid multiple procedures to one site.
- Be gentle, to avoid separating epidermis from dermis.
- Check biopsy site for signs of infection until healing occurs.
- Avoid freezing tissue for electron microscopy because cellular detail will then be destroyed (Table 22.1).
- For specimens undergoing routine microscopic examination, avoid placing biopsy specimen in or on saline because artifactual hydropic degeneration of basal cells and subepidermal bullous formation may occur.



FIG. 22.1. Punch skin biopsy. Top (inset): Disposable biopsy punch. Bottom (inset): Cutting the dermal pedicle.

G. Technique (6,8,17,19,20) See Fig. 22.1.

- Restrain and position patient.
- Choose site for biopsy.
 - For suspected malignant lesions, choose more atypical areas if unable to excise completely.

- For large or chronic lesions, obtain specimen from periphery, including some normal skin.
- For most dermatoses, choose site of early or fully developed, but not end-stage, lesion.
- For acute eruptions and bullous disease, choose an early lesion, including some normal skin.
- For discrete small lesions, try to leave 1- to 2-mm margins of normal skin around the lesions.
- Avoid excoriated, crusted, or traumatized lesions.
- Shave skin, if necessary.
- Prepare as for minor procedure (see Chapter 4).
- Inject 0.25 to 0.5 mL of lidocaine (with/without epinephrine) intradermally beneath the lesion.
- Wait 5 minutes.
- Stretch skin surrounding lesion taut.
- Carefully place punch over the lesion and twist in rotary back-and-forth cutting motion until subcutaneous fat is obtained. Biopsy should include epidermis, full thickness of dermis, and some subcutaneous fat.
- Remove punch.
- Use blunt forceps in one hand to grasp the lateral edge of the biopsy specimen and elevate it, utilizing care to avoid crush artifact.
- Use scalpel blade or scissors in the other hand to cut the punch specimen at its base, as deep into the subcutaneous fat tissue as possible.
- Place specimen in container with appropriate preservative or transport medium.
- Label container with patient name, date, and exact site of biopsy.
- Control bleeding at site of biopsy with gentle pressure on sterile 4- x 4-in gauze square.
- Approximate wound margins and apply Dermabond. No further care is required.
- If suture or Steri-Strips are placed, leave for 5 days on face and for 12 days on trunk, limbs, or scalp.
- Although not recommended by the author, some practitioners allow the wound to heal by secondary intention. If no suture is placed, expect healing by primary epithelialization in 7 to 14 days, with a residual white area a few millimeters in diameter if the biopsy extended to the dermis–subcutaneous fat interface.

TABLE 22.1 Punch Biopsy Preservatives and Transport Media

Transport Medium	Indications	
Formalin 10%	Routine microscopic evaluation	
Michel's Medium or Saline-soaked gauzeBlistering or autoimmune disorders (immunofluorence)		
	Electron microscopy	
	Immunoperoxidase	

H. Complications (6)

- Infection
- Unsightly scarring or keloid formation (rare)
- Excessive bleeding (rare, except in patient with coagulation defect)
- Pathologic uncertainty

References

1. Fretzin D. Biopsy in vesiculobullous disorders. Cutis. 1977; 20:639.

2. Graham J, Barr R. Papulosquamous eruptions: usefulness of biopsy in establishing diagnosis. Cutis. 1977;20:629.

3. Hazelrigg D, Jarratt M. Diagnosis of scabies. South Med J. 1975;68:549.

4. Montes L. How useful is a biopsy in a case of suspected fungal infection? Cutis. 1977;20:665. P.127

5. Roses D, Ackerman A, Harris M, et al. Assessment of biopsy technique and histopathologic

interpretations of primary cutaneous malignant melanoma. Ann Surg. 1979;189:294.

6. Solomon L, Esterly N. Diagnostic procedures. In: Solomon L, Esterly N, eds. Neonatal Dermatology. Philadelphia: Saunders; 1973:29.

7. Soltani K, Pacernick L, Lorincz A. Lupus erythematosus-like lesions in newborn infants. Arch Dermatol. 1974;110:435.

8. Thompson J, Temple W, Lafreniere R, et al. Punch biopsy for diagnosis of pigmented skin lesions. Am Fam Physician. 1988;37:123.

9. Carpenter S, Karpati G, Andermann F. Specific involvement of muscle, nerve and skin in late infantile and juvenile amaurotic idiocy. Neurology. 1972;22:170.

10. Farrell D, Sumi S. Skin punch biopsy in the diagnosis of juvenile neuronal ceroidlipofuscinosis. Arch Neurol. 1977;34:39.

11. Fleisher L, Longhi R, Tallan H, et al. Homocystinuria: investigations of cystathionine synthase in cultured fetal cells and the prenatal determination of genetic status. J Pediatr. 1974;89:677.

 Martin J, Ceuterick C. Morphological study of skin biopsy specimens: a contribution to the diagnosis of metabolic disorders with involvement of the nervous system. J Neurol Neurosurg Psychiatry. 1978;41:232.
Martin J, Jacobs K. Skin biopsy as a contribution to diagnosis in late infantile amaurotic idiocy with curvilinear bodies. Eur Neurol. 1973;10:281.

14. O'Brien J, Bernet J, Veath M, Paa D. Lysosomal storage disorders: diagnosis by ultrastructural examination of skin biopsy specimens. Arch Neurol. 1975;32:592.

15. Spicer S, Garvin A, Wohltmann H, et al. The ultrastructure of the skin in patients with mucopolysaccharidoses. Lab Invest. 1974;31:488.

16. Cooper JT, Goldstein S. Skin biopsy and successful fibroblast culture. Lancet. 1973;2:673.17. Arndt KA. Operative procedures. In: Arndt KA, ed. Manual of Dermatologic Therapeutics. Boston: Little, Brown; 1978: 223.

18. Golbus M, Sagebiel R, Filly R, et al. Prenatal diagnosis of ichthyosiform erythroderma (epidermolytic hyperkeratosis) by fetal skin biopsy. N Engl J Med. 1980;302:93.

19. Ackerman AB. Biopsy: why, where, when, how? J Dermatol Surg. 1975;1:21.

20. Ruiz-Maldonado R, Parish LC, Beare JM. Therapeutic aspects of pediatric dermatology. In: Ruiz-Maldonado R, Parish LC, Beare JM, eds. Textbook of Pediatric Dermatology. Philadelphia: Grune & Stratton; 1989:50.

23-Ophthalmic Specimen Collection

Jennifer A. Dunbar

A. Indications

- To obtain specimen for testing to determine the cause of conjunctivitis (Table 23.1)
 - Neonatal conjunctivitis is considered an ocular emergency (1).
 - Neonatal conjunctivitis is defined as any conjunctivitis presenting in an infant during the first 28 days of life (2).
 - Signs and symptoms include diffuse conjunctival injection with mucoid, purulent, or watery ophthalmic discharge.
 - Conjunctivitis may be the presenting sign of coexisting life-threatening systemic infection.
 - Both bacterial and viral pathogens cause corneal ulceration and opacity, which may lead to blindness. Neisseria gonorrhea or Pseudomonas species may rapidly perforate the globe.
- To test for colonization when a conjunctivitis epidemic is suspected in the neonatal intensive care unit
 - Conjunctivitis is common in the neonatal intensive care unit, affecting up to 6% of infants (3).
 - The eye may be contaminated by respiratory secretions, with coagulase-negative Staphylococcus, S. aureus, and Klebsiella species reported as the most common pathogens. These are associated with ventilator or nasal continuous positive airway pressure (3).
 - Conjunctivitis epidemics have been associated with routine ophthalmic screening in the neonatal intensive care unit (4).
 - Serratia marcessens, Klebsiella species, and Adenovirus epidemics have been described (4,5).
 - Systemic illness and death have been associated with conjunctivitis epidemics in the neonatal intensive care unit (4,5).

Because conjunctivitis of the newborn is often a sign of a maternally transmitted systemic infection, work-up includes maternal history of vaginal symptoms or discharge and maternal vaginal culture results (6).

- B. Relative Contraindications
 - Corneal epithelial defect
 - If fluorescein staining of the cornea reveals an epithelial staining defect, then corneal ulceration may be present. This requires referral to an ophthalmologist.

C. Special Considerations for Ophthalmic Specimen Management

- Conjunctival scrapings are the specimen of choice because many pathogens are intraepithelial (7).
- The ocular specimen size is small; therefore, special care is given to specimen handling.
- Direct placement of the conjunctival scrapings on slides for staining and direct plating onto culture medium at the bedside will maximize the yield.
- Communication with laboratory personnel regarding specimen handling improves culture results (8).

D. Materials

- Equipment for staining the cornea to rule out epithelial defect
 - Fluorescein dye or strips
 - Wood lamp or other blue light source

- Equipment for obtaining specimen
 - Choose topical anesthetic (optional):
 - 0.5% preservative-free tetracaine in unit-dose containers (Alcon Laboratories, Fort Worth, TX, USA)
 - Preservative-free lidocaine (Elkins-Sinn, Cherry Hill, NJ, USA)
 - Cocaine 4% preserved with 0.5% sodium benzoate (Schein Pharmaceuticals, Port Washington, NY, USA) diluted to 1% to 2% with sterile water.

Many topical ophthalmic anesthetics contain preservatives that may inhibit bacterial growth in culture. For this reason, some physicians choose to perform the procedure without anesthetic. Nevertheless, this may be quite painful for the infant. The above-mentioned anesthetics are preservative-free or have been shown not to inhibit bacterial growth (9).

- Sterile cotton swabs may be used to evert the eyelids but are not recommended for specimen collection.
- Choose instrument to obtain cultures:
 - Calcium alginate swabs
 - Sterile Dacron polyester-tipped applicator (Harwood Products Company, Guilford, ME, USA)

Calcium alginate swabs have been shown to yield equal or better organism retrieval in cultures than spatulas or Dacron swabs (10,11). Moistening the swab with tripticase soy (Becton Dickenson and Company, Cockeysville, MD, USA) broth or other culture medium enhances results. However, spatulas have been shown to provide better samples in smear than swabs. Spatulas preserve the conjunctival epithelial cells better, thus providing better opportunity for diagnosing pathogens with intracellular organisms or inclusions (12). Calcium alginate swabs may interfere with immunoassays.

- Choose instrument for scraping the conjunctiva:
 - Kimura Platinum E-109 spatula (Storz Instrument Co., St. Louis, MO, USA)
 - Nasopharyngeal swab with metal handle bent for scraping
 - Calcium alginate swabs

If spatulas are not available, then swabs should be used vigorously on the tarsal conjunctival surface so as to $d\tilde{A}$ [©]bride epithelial cells.

- Equipment for obtaining microscope slides
 - Frosted, etched glass slides
 - Microslide holders
 - Pencil or marker for labeling

Test	Organisms Identified	Finding
Stain		
Gram's stain	Neisseria gonorrhea	Gram-negative diplococci
Giemsa stain	Chlamydia trachomatis	Intraepithelial intracytoplasmic inclusions
Papanicolaou stain	Herpes simplex virus	Multinucleate giant cells and inclusion-bearing cells
Direct antigen detection techniques		-
Immunofluorescent indicator system	Chlamydia trachomatis	
Immunosorbent assay (ELISA)	Chlamydia trachomatis	
	Herpes simplex virus	
Fluorescein-labeled monoclonal antibodies	Chlamydia trachomatis	
(Syva MicroTrack)		
Indirect fluorescence	Herpes simplex virus	
Culture		
Specific media		
Thayer–Martin	Neisseria gonorrhea	
Aerobic	Gram-positive and gramnegative bacteria	
Anaerobic	Anaerobic bacteria	
Viral transport	Herpes simplex virus	
Chlamydia culture (McCoy culture)	Chlamydia trachomatis	
ELISA. enzyme-linked immunosorbent assa	av.	

TABLE 23.1 Analysis of Conjunctival Scrapings

E. Equipment for Identifying Chlamydia

Equipment for nonculture chlamydial studies •

Traditionally, McCoy culture was considered the $\hat{a} \in \hat{a}$ gold standard $\hat{a} \in \hat{a}$ for identification of Chlamydia. However, cultures take several days to provide results, and specimens collected in the first few days of life may have less yield on culture because elementary bodies often take several days to form in neonates (13,14). The nonculture tests listed below, such as direct immunofluorescence, enzyme-linked immunosorbent assay, and polymerase chain reaction (PCR), all perform well for ocular specimens and provide a result more rapidly. Polymerase chain reaction is now commercially available and is probably the most sensitive nonculture test for Chlamydia available (14). Real-time PCR is becoming available for ophthalmic specimens when Chlamydia and/or viral infections such as herpes virus are suspected (15,16). Check with the laboratory involved regarding validation of PCR for ophthalmic specimens.

- For chlamydial DFA stain: Syva MicroTrak Chlamydia trachomatis specimen collection kit (Trinity Biotech, Co., Wicklow, Ireland)
- For chlamydial enzyme-linked immunosorbent assay: Place specimen in media advised by the laboratory performing the study.
- For chlamydial polymerase chain reaction: Place specimen in transport medium appropriate 0 for the assay used (13). An example is M4 medium for the transport of viruses and Chlamydia (Remel, Lenexa, KS, USA).
• Media for Chlamydia culture

Specimens should be plated onto culture medium at the bedside. Each laboratory will have specific media available for a particular type of organism. Check with the laboratory involved for the appropriate medium for that hospital. The following list is a suggestion of classic media used for each type of organism.

- Bacterial culture media
 - Trypticase soy broth
 - Blood agar plate
 - Chocolate agar plate for Hemophilus influenzae, Neisseria gonorrhea
 - Thayer-Martin medium if gonorrhea suspected
- Virus-holding medium, i.e., M4 medium for the transport of viruses and Chlamydia (Remel, Lenexa, KS, USA)
- Chlamydia culture transport medium, i.e., M4 medium for the transport of viruses and Chlamydia (Remel, Lenexa, KS, USA)
- Sabourad's agar if fungal conjunctivitis suspected

F. Technique

- Method for staining the cornea for epithelial defect
 - Instill a very small amount of fluorescein in the lower conjunctival fornix by lightly touching the tear film with a fluorescein strip. Flooding the eye with fluorescein may obscure a small corneal epithelial defect.
 - Evaluate the cornea for staining with a Wood lamp or other blue light source.
 - If a corneal epithelial defect is present, the cornea may be infected and an ophthalmologist should be consulted.
 - Herpes virus may present in the neonate as a geographic-shaped epithelial effect rather than a dendrite.



FIG. 23.1. Everting the upper eyelid.

- Method for everting eyelids
 - Upper lid (Fig. 23.1) 0
 - Grasp lashes and border of lid between thumb and index finger of nondominant hand.
 - Draw lid downward and away from eyeball.
 - Indent upper lid, with handle of cotton-tipped applicator held in dominant hand and pull lid back and upward over applicator.
 - Remove applicator and hold lid in place with nondominant hand by gently pressing border of lid against superior orbital margin.
 - Lower lid (Fig. 23.2) 0
 - Place index finger of nondominant hand on margin of lower lid.
 - Pull downward. •
- Method for obtaining cultures

Obtain cultures prior to conjunctival scraping. Take separate cultures from each eye with a separate sterile swab for each type of medium desired. Culture and label each eye separately, even if only one eye is symptomatic. The uninfected eye can serve as a control for indigenous flora (8).



FIG. 23.2. Using Kimura platinum spatula to take scraping from lower eyelid.

- Moisten calcium alginate swabs with trypticase soy broth or other liquid culture medium.
- Evert eyelid.
- Apply swab to bulbar and palpebral conjunctiva of upper and lower fornices of eye.
- Apply swab directly to culture medium plates at the bedside with a single row of C-shaped inoculation streaks. Monitoring the growth of organisms along the shape of the inoculation streaks may help the laboratory in the diagnosis of the cultured pathogen.
- Use a separate sterile swab for each culture plate or culture vial.
- Label cultures meticulously with eye cultured (right or left) and part of eye cultured (conjunctiva, lid margin, etc.).
- Incubate cultures immediately.
- Method for obtaining conjunctival scrapings for smear and nonculture Chlamydia tests
 - Evert eyelid as described above.
 - Instill topical anesthetic into conjunctival fornix, if desired.
 - Swab off excess discharge.
 - Take scraping 2 mm from eye margin. (Normal keratinized epithelium from the lid margin may confound results of smear.)
 - Pass spatula two to three times in the same direction, avoiding bleeding.
 - Spread specimen from spatula gently into a monolayer on a clean glass slide and label.
 - \circ $\;$ Fix smears as required for proposed smears and nonculture Chlamydia tests.
 - Repeat with separate sterile spatula on second eye.

G. Interpretation of Conjunctival Cytology

- Cellular reaction
 - Polymorphonuclear reaction
 - Bacterial infections
 - Chlamydial infection
 - Very severe viral infection

- Mononuclear reactions: viral infection
- Eosinophilia and basophilia: allergic states
- Plasma cells: chlamydial infection
- Intraepithelial cell inclusions
 - Chlamydial infection
 - Acidophilic inclusions in cytoplasm, capping epithelial cell nuclei
 - Basophilic "initial bodies†in cytoplasm
 - Viral infection

Giant, multinucleated epithelial cells may be seen (e.g., herpetic keratoconjunctivitis).

H. Complications of Scraping

- Conjunctival bleeding
 - Mild conjunctival bleeding, usually self-limiting, frequently occurs.
 - Instill erythromycin ophthalmic ointment.
- Corneal injury
 - Keep the spatula blade flat against the tarsal conjunctiva at all times to avoid trauma to the cornea.
 - Corneal injury is confirmed by a staining defect on fluorescein staining.
 - If corneal injury occurs, instill erythromycin ophthalmic ointment and contact an ophthalmologist.
- Transfer of infection from infected to noninfected eye

This complication is avoided by using separate sterile instruments when taking samples from each eye.

• Ocular irritation, pain, photophobia, lacrimation, swelling, and hyperemia

These problems are usually mild and self-limited.

References

1. De Toledo AR, Chandler JW. Conjunctivitis of the newborn. Infect Dis Clin North Am. 1992;6:807–813.

2. World Health Organization. Conjunctivitis of the Newborn: Prevention and Treatment at the Primary Health Care Level. Geneva: World Health Organization; 1986:1–23.

3. Haas J, Larson E, Ross B, et al. Epidemiology and diagnosis of hospital acquired conjunctivitis among neonatal intensive care unit patients. Pediatr Infect Dis J. 2005;24(7):586–589.

4. Faden H, Wynn RJ, Campagna L, et al. Outbreak of adenovirus type 30 in the neonatal intensive care unit. J Pediatr. 2005;146(4):523–527.

5. Casolari C, Pecorari M, Fabio G, et al. A simultaneous outbreak of Serratia marcescens and Klebsiella pneumoniae in a neonatal intensive care unit. J Hosp Infect. 2005;61(4): 312–320.

6. Hammerschlag MR. Neonatal conjunctivitis. Pediatr Ann. 1993;22:326–361.

7. O'Hara MA. Ophthalmia neonatorum. Pediatr Clin North Am. 1993;40:715–725.

8. Miller JM, ed. A Guide to Specimen Management in Clinical Microbiology. 2nd ed. Washington, DC: American Society for Microbiology Press; 1999.

9. Mullin GS, Rubinfeld RS. The antibacterial activity of topical anesthetics. Cornea. 1997;16:662–665.
10. Benson WH, Lanier JD. Comparison of techniques for culturing corneal ulcers. Ophthalmology. 1992;99:800–804.

11. Jacob P, Gopinathan U, Sharma S, et al. Calcium alginate swab versus Bard Parker blade in the diagnosis of microbial keratitis. Cornea. 1995;14:360–364.

12. Rapoza PA, Johnson S, Taylor HR. Platinum spatula vs Dacron swab in the preparation of conjunctival smears [Letter]. Am J Ophthalmol. 1986;102:400–412.

13. Talley AR, Garcia-Ferrer F, Laycock KA, et al. Comparative diagnosis of neonatal chlamydial conjunctivitis by polymerase chain reaction and McCoy cell culture. Am J Ophthalmol. 1994;117: 50–57. 14. Hammerschlag MR, Roblin PM, Gelling M, et al. Use of polymerase chain reaction for the detection of Chlamydia trachomatis in ocular and nasopharyngeal specimens from infants with conjunctivitis. Pediatr Infect Dis J. 1997;16:293–297.

15. Kowalski RP, Thompson PP, Kinchington PR, et al. Evaluation of the Smart Cycler II system for realtime detection of viruses and Chlamydia from ocular specimens. Arch Ophthalmol. 2006;124(8): 1135–1139.

16. Chichili GR, Athmanathan S, Farhatullah S, et al. Multiplex polymerase chain reaction for the detection of herpes simplex virus, varicella-zoster virus and cytomegalovirus in ocular specimens. Curr Eye Res. 2003;27(2):85–90.

24-Perimortem Sampling

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The death of a newborn is devastating for parents and extended families. This is particularly true when the cause of death is a suspected but unconfirmed genetic or metabolic disorder or when the cause of death is unknown. Approximately one third of pediatric admissions have a genetic component to their illness (1), and genetic disease is a major contributing factor in 15% to 25% of all infant deaths (2). Additionally, more than 400 inborn errors of metabolism have been described, and although treatment may be available, many infants die before the diagnosis is made or even considered (3). The majority of inborn errors are autosomal recessive, implying a 25% recurrence risk for future pregnancies. For many of these disorders, arriving at a diagnosis often hinges on timely perinatal and postnatal sampling. Perhaps the greatest service a neonatologist can render to the family of a dying child is appropriate evaluation and timely collection of samples for diagnosis.

During the discussion with the family to withdraw life support or as soon after the infant expires as possible, discuss with the parents the importance of collecting perinatal fluid and tissue samples and performing an autopsy. Postmortem examinations can be crucial in determining cause of death and may have implications for the health of other family members. Such examinations have been shown to alter the recurrence risks for future pregnancies in 22% to 26% of cases (4,5,6 and 7).

If family members are resistant to a complete autopsy, then a limited study with exclusions or permission to obtain tissue samples can also be extremely helpful. Noninvasive techniques such as postmortem magnetic resonance imaging (MRI), especially in conjunction with tissue sampling, may be extremely helpful in arriving at a diagnosis and may be more acceptable to the family (8).

A. Importance of Perimortem Diagnostic Tests (3,9)

- Establish the correct diagnosis.
- Give parents an accurate explanation of the cause of death and allay parental fears about what did not cause the child's death.
- Permit appropriate testing of asymptomatic parents and siblings, potentially facilitating prenatal diagnosis and preventing mortality and morbidity.
- Provide genetic counseling for recurrence risks and discussion of reproductive options and prenatal diagnosis.
- Dispel allegations of abuse or neglect.

B. Indications (10 11,12,13 and 14)

- Suspected inborn error of metabolism
- Suspected genetic disease or chromosome abnormality
- Suspected infectious disease
- Unexplained death

C. Precautions

- Informed consent is essential for all procedures.
- To prevent bacterial contamination, use sterile procedures even for postmortem sampling.

D. Perimortem Evaluation

- Timely consultation with a clinical and/or biochemical geneticist as soon as a genetic or metabolic disorder is suspected is key in evaluating the neonate and guiding the correct retrieval, collection, and storage of appropriate samples for analysis. When a genetic or biochemical disorder is suspected, the perimortem collection of samples should be considered an emergency procedure (3).
- Careful family history including three-generation pedigree. Pay special attention to neonatal deaths or stillbirths, pregnancy losses, and sudden or unexplained infant deaths. Ask about consanguinity, ethnicity, and obtain an obstetric history. Based on the clinical findings, consider work-up for a thrombophilia disorder in the mother (3).
- Dysmorphology examination (performed by a clinical geneticist with expertise in dysmorphology if possible)
 - Document both negative and positive findings.
 - Pay special attention to:
 - Growth parameters (large or small for gestational age)
 - Distribution, abundance, and texture of the hair
 - Subtle facial variations including ear configuration and placement and presence of facial or palatal clefts
 - Eyes for spacing, palpebral fissure length and slant, eyelid malformation, colobomas, microphthalmia, cataracts, or obvious retinal changes
 - Chest size, sternal or rib malformations, abdominal wall or spinal defects
 - Digital abnormalities including syndactyly, polydactyly, or positioning defects
 - Genital or rectal anomalies
- Photographs
 - Obtain photographs of as high a quality as possible.
 - A fuzzy Polaroid is better than no photo at all.
 - Obtain both anteroposterior and lateral views.
 - Obtain close-up photographs of any abnormalities.
 - Try to obtain photographs without obscuring tubes or tape.
- Radiographs
 - If a skeletal disorder is suspected, radiographs of the infant are of supreme importance and often diagnostic (15).
 - For small infants, a "babygram†or Faxitron (Faxitron X-ray Corp., Wheeling, IL, USA) study can be used.
 - Anteroposterior and lateral views should be obtained, as well as hands and feet and any extremity variations.
- Magnetic resonance imaging (MR-autopsy) (8)
 - o Advantages
 - MR is noninvasive and may be acceptable to parents who refuse full autopsy for fear of disfigurement.

- High-quality images can be obtained especially of the central nervous system (CNS anomalies are present in 1% of all births) (16), which is often difficult to evaluate by conventional autopsy.
- The organs are examined in situ; therefore, complex malformations involving several organs can be evaluated in correct anatomic position. This can be particularly important for CNS and spinal cord anomalies.
- MR-autopsy can be performed up to 72 hours after death.
- o Disadvantages
 - No tissue samples can be obtained for culture, chromosome, or metabolic analysis.
 - Complex cardiac malformations and anomalous connections between hollow organs (i.e., tracheo-esophageal fistula or bowel perforations) are difficult to detect by MR-autopsy.
 - Skeletal malformations may be missed.
 - Most hospital MR staff are not familiar with examining deceased neonates and will need to be educated about the value of such studies.

E. Perimortem Sampling (3,10)

- Blood samples
 - Contact the hospital laboratory immediately to save any unused portions of blood specimens sent earlier for other routine tests. Have the serum or plasma separated and frozen and any unused urine or cerebrospinal fluid frozen.
 - Contact the state laboratory to retrieve any unused portion of blood spots sent for newborn screening. If not previously reported, inquire about the results of the newborn screening.
 - When death is imminent or the family has elected to withdraw life support measures for an infant, obtain blood samples for further analysis (Table 24.1).
 - Dried blood spots on filter paper (at least two or three newborn screening cards) stored at room temperature but not in a plastic bag.
 - Whole blood
 - 5 mL in lithium heparin tube (separated within 20 minutes of collection and stored at -80ŰC)
 - 5 mL in EDTA tube for DNA extraction (can be stored for 48 hours at 4°C prior to extraction)
 - 5 mL in sodium heparin (can be stored at room temperature or at 4°C overnight for chromosome analysis; do not freeze)
- Urine samples (10)
 - May be used for urine organic acid and amino acid analysis
 - Collect by catheterization, Crede's method, or bladder tap and freeze immediately. If the sample is contaminated with blood, centrifuge the sample to remove red cells, then freeze. Washing out the bladder with a small amount of sterile saline may yield enough sample for organic acid analysis.
 - $\circ~$ Volume of 5 to 10 mL is ideal, but even the smallest amount of urine available may be useful. As little as 100 ŵL can be sufficient for organic acid analysis by gas chromatography-mass spectrometry
- Hair samples
 - Reserve a few strands of hair (plucked from the scalp) for microscopic analysis (17)

TABLE 24.1 Dioou Sample Conection			
Laboratory Test	Volume	Tube	Handling
	(mL)		
Karyotype	1.5	Sodium	Room temperature or refrigerate, do not
		heparin	freeze
Chromosome microarray	3	Sodium	Room
analysis temperature	3	heparin	
		K ₃ EDTA	
Plasma amino acids	1–2	Lithium	On ice; if not processed immediately,
		heparin	separate plasma and freeze
Plasma carnitine	1–3	Lithium	On ice
		heparin	
Very-long-chain fatty acids	3	$K_3 EDTA$	Room temperature
White blood cells for	3	K ₃ EDTA	Room temperature
enzymes/DNA			-

TADLE 24.1 Blood Somple Collection

EDTA, ethylenediaminetetraacetate.

- Cerebrospinal fluid
 - For suspected metabolic disorder, specimen should be obtained prior to death and immediately frozen at -80 \hat{A} °C (10,18).
 - A postmortem sample of cerebrospinal fluid may be obtained by passing a needle through the anterior fontanelle using aseptic technique and may be useful for detecting an infectious disorder (4).
- Skin biopsy (19,20)
 - Skin biopsy specimens may be subjected to microscopic examination or used for culturing skin fibroblasts. Skin fibroblasts can be used for chromosome analysis, for DNA isolation for mutation detection, and for enzyme analysis. Specimens should be obtained as close to the time of death as possible, but skin biopsy up to 2 to 3 days postmortem may still produce a viable culture. In this case, several small biopsies taken from a number of separate sites (skin or fascia) may be more successful.
 - Clean the inside of the forearm or the anterior thigh with numerous alcohol swabs.
 - Povidone–iodine should not be used because it can impair cell growth.
 - Use a 3- or 5-mm punch biopsy or a scalpel to obtain a full thickness of skin. A small piece of sterile fascia (10) or a fine-needle aspiration of tissue (21) may also be used for fibroblast culture.
 - Place the biopsy tissue in a culture medium such as Hanks' or minimal essential medium (MEM), or a viral transport medium. If these are unavailable, place in sterile saline.
 - Refrigerate specimen or transport at room temperature. Do not freeze.
 - Cells can be cultured and then archived in liquid nitrogen for decades and still be successfully recovered for analysis.
- Muscle biopsy (10,22,23)
 - If a neonatal muscular dystrophy or a mitochondrial disorder of energy metabolism is suspected, try to obtain a skeletal muscle biopsy as close to the time of death as possible, but within 2 to 4 hours. A percutaneous or incisional muscle biopsy can be performed at the bedside. A neurologist or surgeon may be more experienced in obtaining an appropriate specimen and/or for providing the muscle clamps necessary to appropriately obtain the specimen.
 - Incisional biopsy. Make a 2- to 3-cm incision over the quadriceps muscle to expose it. Muscle clamps should be applied prior to removal of the muscle fibers in order to prevent the muscle from contracting. Three muscle fibers should be removed and processed as follows.

- Place one on a saline-soaked Telfa pad (Kendall Company, Mansfield, MA, USA) and refrigerate. Do not have the muscle floating in saline. This will be used for light microscopy.
- Place a second specimen in 1.5% glutaraldehyde/1% formaldehyde in 0.12 M Sorensen's buffer and place in the refrigerator. If this is not available, place the sample in formalin and leave at room temperature. If neither of these is available, place the specimen on a saline-soaked Telfa pad and refrigerate. This will be used for electron microscopy.
- Wrap a third specimen in aluminum foil and snap-freeze in isopentane if available. Alternatively, the specimen can be snap-frozen in liquid nitrogen or dry ice and stored at - 80ŰC. This will be used for enzyme analysis.
- Percutaneous biopsy. If an incisional biopsy is not possible, three cores of quadriceps muscle should be obtained percutaneously using a biopsy needle and processed as described above.
- Liver biopsy (11,24,25)
 - If there is hepatic failure or an inborn error of metabolism is suspected, try to obtain a biopsy of the liver as close to the time of death as possible. If the autopsy will not be performed for several hours and the hospital pathologist concurs, or if the family has declined autopsy but permitted tissues to be obtained, an open wedge biopsy or percutaneous biopsy can be performed at the bedside. If the biopsy has to be delayed for any reason, specimens obtained for biochemical analysis even up to 72 hours after the time of death may provide significant results (11).
 - Wedge biopsy. Make a 2-cm skin incision just below the right costal margin. With forceps, lift up the right lobe of the liver, and with a clean scalpel, remove a wedge. Cut the wedge into several 5-mm cubes and place in a plastic cryotube. Snap-freeze the tube in liquid nitrogen, or if this is not available, bury it in dry ice. Store at 280ŰC
 - Percutaneous biopsy. A 16- or 18-gauge Menghini needle (Allegiance Healthcare Corp., 0 McGraw Park, IL, USA), a Jamshidi needle (Pharmaseal; Baxter Healthcare Corp., Valencia, CA, USA), a 16-gauge Klatskin needle (Becton Dickinson, Franklin Lakes, NJ, USA), a Tru-Cut needle (Travenol Laboratories, Deerfield, IL, USA), or a spring-loaded disposable liver biopsy needle (Monopty; Bard Peripheral Technologies, Covington, GA, USA) may be used. With the infant positioned supine, mark a site slightly below the right midaxillary line, at the ninth or tenth rib. Make an incision in the skin 1/8 in long with a no. 11 scalpel blade above the rib. Attach a locking 10-mL syringe filled with 5 mL of normal saline to the needle. Flush to remove the air and direct the liver biopsy needle through the skin incision, parallel to the surface the infant is on, directed toward the xiphoid, nipple, or opposite shoulder. Flush with 1 mL of saline to clear the cannula. Advance 2 to 3 cm while creating negative pressure in the syringe by pulling back the plunger 1 to 2 mL and turning it into the locking position. Withdraw the needle and syringe assembly. If the automatic spring-loaded system is used, negative pressure is not required. Several cores of liver should be obtained using a biopsy needle and processed as described above.
- Placenta and umbilical cord
 - Ensure that the placenta is sent to the pathologist for evaluation for every sick infant (26).
 - Examination of the placenta may provide clues to maternal or fetal vascular problems (27), in utero infectious or abnormal inflammatory responses (28), or lysosomal storage disorders (29).
 - A sample of the umbilical stump in the newborn may identify invasive inflammation in cases of sepsis.

F. The Postmortem Professional and Family Conference (8,30)

- It is important to keep in contact with the family. Apprise them of what studies are to be performed on their deceased infant and how long they will take, and what samples remained stored if further diagnostic testing becomes available.
- If not done previously, obtain a complete three-generation family history as described in D.2. Obtain the services of a geneticist or genetic counselor in reviewing the family history and assigning recurrence risks.
- Review the prenatal history for any exposures to infection or potential teratogens including environmental toxins. Remember, the parents are looking for answers and will usually be eager to share any bit of information they think might be helpful in arriving at a cause of death.
- If possible, schedule a multidisciplinary conference involving the neonatologists as well as the clinical or biochemical geneticist, pathologist, radiologist, or other subspecialists who cared for the child or who have taken part in the postmortem evaluation. Carefully review the findings and arrive at a diagnosis or possible diagnoses, including recurrence risks and reproductive options available to the couple. Identify any family members who may be asymptomatic carriers or at risk for the disorder so that additional testing can be offered.
- When the autopsy and any special studies are complete, schedule a conference with the family. Review the findings. It is often important to explain what did not lead to the infant's death as well as the suspected or confirmed final diagnosis. With the help of a geneticist or genetic counselor, discuss the recurrence risks for future pregnancies and what types of studies (e.g., chromosome or biochemical analysis, ultrasound, MRI, or fetal echocardiography) could be used to monitor subsequent pregnancies. If available, offer testing to family members who are suspected carriers or who may be asymptomatic but at risk for the disorder (e.g., the deceased child's siblings).
- Provide the family and the referring physician with a copy of the autopsy report and any diagnostic testing results for their records. Provide a letter summarizing the postmortem conference. These may become very important to them in the future.
- Finally, remain available to the family to answer any additional questions or concerns that may arise over the next several weeks to months as they grieve and come to accept of the loss of their child.

References

1. Weatherall DJ. The New Genetics and Clinical Practice. Oxford: Oxford University Press; 1991:32–37.

2. Penchaszadeh VB. Reproductive genetic testing from an international perspective: impact on women in underdeveloped countries. Fetal Diagn Ther. 1993;8:202–209.

3. Christodoulou J, Wilcken B. Perimortem laboratory investigation of genetic metabolic disorders. Semin Neonatol. 2004;9:275–280.

4. Bove KE. Practice guidelines for autopsy pathology. The perinatal and pediatric autopsy. Arch Pathol Lab Med. 1997;121:368–376.

5. Faye-Peterson OM, Guinn DA, Wenstrom KD. Value of perinatal autopsy. Obstet Gynecol. 1999;94:915–920.

6. Rushton DI. Prognostic role of the perinatal postmortem. Br J Hosp Med. 1994;52:450–454.

7. Doyle LW. Effects of perinatal necropsy on counseling. Lancet. 2000;355:2093.

8. Huisman TAGM. Magnetic resonance imaging: an alternative to autopsy in neonatal death? Semin Neonatol. 2004;9:347–353.

9. Bennett MJ, Rinaldo P. The metabolic autopsy comes of age. Clin Chem. 2001;7:1145–1146.

10. Olpin SE. The metabolic investigation of sudden infant death. Ann Clin Biochem. 2004;41:282–293.

11. Rinaldo P, Yoon H, Yu C, et al. Sudden and unexpected neonatal death: a protocol for postmortem diagnosis of fatty acid oxidation disorders. Semin Perinatol. 1999;23:204–210.

12. Chace DH, Diperna JC, Mitchell BL, et al. Electrospray tandem mass spectrometry for acylcarnitines in dried postmortem blood specimens collected at autopsy from infants with unexplained cause of death. Clin Chem. 2001;47:1166–1182.

13. Anderson C, Ramsay JA, Nogee LM, et al. Recurrent familial neonatal deaths: hereditary surfactant protein B deficiency. Am J Perinatol. 2000;17:219–224.

14. Van den Veyver IB, Ni J, Bowles N, et al. Detection of intrauterine viral infection using the polymerase chain reaction. Mol Genet Metab. 1998;63:85–95.

15. Foote GA, Wilson AJ, Stewart JH. Perinatal postmortem radiographyâ€"experience with 2500 cases. Br J Radiol. 1978;51:351â€"356.

16. Pinar H, Tatevosyants N, Singer DB. Central nervous system malformations in a perinatal/neonatal autopsy series. Pediatr Dev Pathol. 1998;1:42–48.

17. Kodama H, Murata Y, Kobayashi M. Clinical manifestations and treatment of Menkes disease and its variants. Pediatr Int. 1999;41:423–429.

18. Hoffman GF, Surtees RA, Wevers RA. Cerebrospinal fluid investigations for neurometabolic disorders. Neuropediatrics. 1998;29:59–71.

19. Lesca G, Boggio D, Bellec V, et al. Trisomy 17 mosaicism in amniotic fluid cells not found at birth in blood but present in skin fibroblasts. Prenat Diagn. 1999;19:263–265.

20. Lundemose JB, Kolvraa S, Gregerson N, et al. Fatty acid oxidation disorders as a primary cause of sudden and unexpected death in infants and young children: an investigation performed on cultured fibroblasts from 79 children who died aged between $0\hat{a}\in$ 4 years. Mol Pathol. 1997;50:212 $\hat{a}\in$ 217.

21. Kurtycz DF, Logrono R, Harris C, et al. Use of fine needle aspiration for fibroblast culture. Pediatr Pathol Lab Med. 1998;18:35–39.

22. Darras BT, Jones HR. Diagnosis of pediatric neuromuscular disorders in the era of DNA analysis. Pediatr Neurol. 2000;23:289–300.

23. Kelly NA, Thomas C. Pathologic quiz case: male infant with generalized hypotonia and absence of respirations at birth. Arch Pathol Lab Med. 2001;125:575–576.

24. Chow CW, Thorburn DR. Morphological correlates of mitochondrial dysfunction in children. Hum Reprod. 2000;15:68–78.

25. Boles RG, Buck EA, Blitzer MG, et al. Retrospective biochemical screening of fatty acid oxidation disorders in postmortem livers of 418 cases of sudden death in the first year of life. J Pediatr. 1998;132:924–933.

26. Altshuler G. Role of the placenta in perinatal pathology (revisited). Pediatr Pathol Lab Med. 1996;16:207–233.

27. Khong TY. Placental vascular development and neonatal outcome. Semin Neonatol. 2004;9:255–263.
28. Redline RW. Placental inflammation. Semin Neonatol. 2004;9:265–274.

29. Roberts DJ, Ampola MG, Lage JM. Diagnosis of unsuspected fetal metabolic storage disease by routine placental examination. Pediatr Pathol. 1991;11:647–656.

30. McHaffie HE, Laing IA, Lloyd DJ. Follow up care of bereaved parents after treatment withdrawal from newborns. Arch Dis Child Fetal Neonat Ed. 2001;84:F125–F128.

25-Abdominal Paracentesis

A. Alfred Chahine

A. Indications

- Therapeutic
 - Massive ascites with cardiorespiratory compromise
- Diagnostic
 - Necrotizing enterocolitis with suspicion of gangrene or perforation, looking for fecal matter or bacteria and white blood cells on a smear (1,2 and 3)
 - \circ Chylous ascites: testing for lymphocytes on cell count of the fluid (4)
 - Urinary ascites: test for creatinine content (5,6)
 - Meconium peritonitis: gross appearance of ascites (7)
 - Biliary ascites: test for bilirubin level.
 - Congenital infectionsâ€"cytomegalovirus (CMV), tuberculosis, toxoplasmosis, syphilis: test for inclusion bodies, treponemes (4,8)
 - Inborn errors of metabolismâ€"sialic acid storage disorders: test for vacuolated lymphocytes and free sialic acid (9,10)
 - Iatrogenic ascites from extravasation of fluid from central venous catheters: test for glucose content

B. Contraindications

Coagulopathy is a relative contraindication; the procedure may be performed with concomitant treatment of thrombocytopenia or coagulopathy.

C. Equipment

- 24- or 25-gauge catheter over a needle (e.g., Angiocath)
- 5- or 10-mL syringe
- Skin topical disinfectant
- Sterile towels
- Tubes for culture, Gram stain, and cell count
- Tuberculin syringe
- Lidocaine (1%)

D. Technique (See Procedures DVD for Video)

- A soft support ("bumpâ€) is placed under the neonate's left flank, to allow as much of the fluid to drain into and allow the intestines to float away from the right lower quadrant (Fig. 25.1).
- The right lower quadrant is prepared with the disinfecting solution.
- A point between the umbilicus and the anterior superior iliac spine one third of the way from the anterior superior iliac spine is chosen.
- The skin, muscles, and peritoneum are infiltrated with the local anesthetic using the tuberculin syringe.
- The 10-mL syringe is connected to the 24-gauge catheter and needle.
- The catheter is pointed toward the back and held at a 45-degree angle (Fig. 25.2).
- The catheter and needle are slowly pushed through the skin, muscles, and peritoneal surface while applying gentle suction on the syringe plunger.
- When peritoneal fluid is aspirated, the catheter is advanced and the needle withdrawn.

- The syringe is connected to the catheter and suction applied to aspirate as much fluid as possible.
- If fluid is not free-flowing, the catheter might be inside the lumen of a piece of intestine or in the retroperitoneum. The catheter is withdrawn and the maneuver repeated with the catheter at a slightly different angle.
- When the fluid stops flowing, the catheter is withdrawn.
- The fluid is distributed into the various tubes and cups for the appropriate studies.
- A bandage is applied.

E. Complications

- Bleeding from the liver or intra-abdominal vessels; may be severe enough to require a laparotomy
- Intestinal perforation; usually inconsequential because the catheter and needle are of small diameter.



FIG. 25.1. Appropriate position and disinfection of abdomen prior to performing paracentesis in preterm neonate.



FIG. 25.2. Entry site and direction of needle for abdominal paracentesis in preterm neonate.

References

1. Ricketts RR. The role of paracentesis in the management of infants with necrotizing enterocolitis. Am Surg. 1986;52(2):61–65.

2. Ricketts RR, Jerles ML. Neonatal necrotizing enterocolitis: experience with 100 consecutive surgical patients. World J Surg. 1990;14:600–605.

3. Kosloske AM. Indications for operation in necrotizing enterocolitis revisited. J Pediatr Surg. 1994;29(5):663–666.

4. Lee YY, Soong WJ, Lee YS, Hwang B. Total parenteral nutrition as a primary therapeutic modality for congenital chylous ascites: report of one case. Acta Paediatr Taiwan. 2002;43(4):214–216.

5. Oei J, Garvey PA, Rosenberg AR. The diagnosis and management of neonatal urinary ascites. J Paediatr Child Health. 2001;37(5):513–515.

6. Ku JH, Kim ME, Jeon YS, et al. Urinary ascites and anuria caused by bilateral fungal balls in a premature infant. Arch Dis Child Fetal Neonatal Ed. 2004;89(1):F92–F93.

7. Shyu MK, Shih JC, Lee CN, et al. Correlation of prenatal ultrasound and postnatal outcome in meconium peritonitis. Fetal Diagn Ther. 2003;18(4):255–261.

8. Nicol KK, Geisinger KR. Congenital toxoplasmosis: diagnosis by exfoliative cytology. Diagn Cytopathol. 1998;18:357–361.

9. Sergi C, Beedgen B, Kopiz J, et al. Refractory congenital ascites as a manifestation of neonatal sialidosis: clinical, biochemical and morphological studies in a newborn Syrian male infant. Am J Perinatol. 1999;16:133–141.

10. Lemyre E, Russo P, Melancon SB, et al. Clinical spectrum of infantile free sialic acid storage disease. Am J Med Genet. 1999;82:385–391.

26-Peripheral Intravenous Line Placement

Leah Greenspan-Hodor Khodayar Rais-Bahrami Martin R. Eichelberger

Percutaneous Method

A. Indications

• Administration of intravenous (IV) medications, fluids, or parenteral nutrition when utilization of the gastrointestinal tract is not possible

B. Equipment

Since the late 1960s, the variety of equipment available for peripheral vascular access has grown from a few sizes of metallic needles and stiff polyethylene tubes to an array of plastic cannulas, single- and multilumen catheters of different sizes and materials, and totally implantable devices (ports). The safest and more effective vascular access is obtained by carefully matching the neonate's size, therapeutic needs, and duration of required treatment with the most appropriate device and technique. Placement of peripheral IV lines is described in this chapter. Placement of central venous lines (excluding ports, which are not used routinely in neonates) is described in Chapter 31. Sterile (Fig. 26.1)

- Povidone iodine swabs or 70% alcohol swabs (or other antiseptic; see Chapter 4)
- Appropriate needle (minimum 24 gauge for blood transfusion)
 21- to 24-gauge IV catheter (preferably shielded)
- Connection for cannula (i.e., T connector)
- 2- x 2-in gauze squares
- Isotonic saline in 3-mL syringe
- Heparinized flush solution (heparin 0.5 to 1 U/mL/mL normal saline) for heparin lock
- Scissors (for scalp IV placement)

Nonsterile

- Tourniquet
- Procedure light
- Materials for restraint (see Chapter 3)
- Transilluminator (optional)
- Warm compress to warm limb if necessary (heel warmer)
- Size-appropriate arm board
- Cotton balls
- Scissors
- Roll of 0.5- to 1-in porous adhesive tape, transparent tape, or semipermeable transparent dressings (1,2,3,4 and 5)
 - If using tape, use the minimum amount necessary on fragile premature skin, and consider using a pectin barrier (Duoderm, ConvaTec/Bristol-Myers Squibb, Princeton, NJ, USA; Hollihesive, Hollister, Libertyville, IL, USA).
 - Transparent tape or dressing will facilitate observation of IV site (Tegaderm, 3M Health Care, St. Paul, MN, USA).
 - Precut self-adhesive taping devices are available from Veni-Gard Jr. (ConMed IV Site Care Products, Utica, NY, USA).

• Pacifier if appropriate. Sucking releases endorphins, which decrease pain. Consider tightly swaddling baby, leaving the limb needed for IV placement exposed. Swaddling is also a comfort measure. Some critically ill infants, such as a baby with persistent pulmonary hypertension (PPHN), may require pain medication, sedation, and/or paralysis prior to any invasive procedure, including IV line placement.

C. Precautions

- Avoid areas adjacent to superficial skin loss or infection.
- Avoid vessels across joints, because immobilization is more difficult.
- Take care to differentiate veins from arteries.
 - Palpate for arterial pulsation.
 - Note effect of vessel occlusion.
 - Limb vessel: arteries collapse, veins fill
 - Scalp vessel: arteries fill from below, veins fill from above



FIG. 26.1. Sterile equipment necessary for peripheral IV line placement.

- Note color of blood obtained (arterial blood is bright red and venous blood is darker).
- \circ Look for blanching of skin over vessel when fluid is infused (arterial spasm).
- If limb requires warming prior to procedure, use a heel warmer (WarmGel, Prism Technologies, San Antonio, TX, USA). "Home-made†compresses such as a diaper soaked in hot water can cause severe thermal injury.
- Cut scalp hair using small scissors to allow for stabilization of the IV (do not shave the area).
- Apply tourniquet correctly (see D5 and Chapter 13).
 - Minimize time applied.
 - Avoid use in areas with compromised circulation.
 - Avoid use for scalp vesselsâ€"tends to increase vessel fragility.
- When using peripheral scalp veins, avoid sites outside the hairline.
- Be alert for signs of phlebitis or infiltration.

- Inspect site hourly.
- Discontinue IV immediately at any sign of local inflammation or cannula malfunction.
- Long plastic catheters are not recommended for use in neonates because their relative rigidity increases the risk of damage to the vascular endothelium, thus increasing the possibility of venous thrombosis (6).
- Arrange tape dressing at IV site to allow adequate inspection or use transparent sterile dressing over site of skin entry (7).

Leibovici (8) was unable to show a positive effect of a daily change of the dressing, as compared with change every 72 hours, on the incidence of infusion phlebitis. Maki and Ringer (3) recommended not removing the transparent dressing until the catheter/ needle is removed.

- Consider use of protective skin preparation in small premature infants to prevent skin trauma upon removal of tape or dressing. No Sting Skin Protectant (3M Health Care, St. Paul, MN, USA) is a non-alcohol-containing product that is available commercially; however, it, as well as other commercially available skin protectants, has not been tested on neonates.
 - Forms tough, protective coating, which bonds to skin
 - Does not require removal when changing dressing
- The use of tincture of benzoin and other products to increase the adherence of tape should be limited, especially on the premature infant. These products create a tighter bond between the tape and the epidermis than the bond between the epidermis and the underlying dermis. This then causes stripping of the epidermis when the tape is removed. Using a protective skin preparation, for example, No Sting Skin Protectant, prior to the application of these products may decrease damage to the skin when tape is removed (9).
- Write date, time, and needle/cannula size on piece of tape secured to site.
- Loop IV tubing and tape onto extremity to take tension off the IV device.
- Limit to two to three placement attempts per person. Monitor carefully for clinical decompensation, particularly the very premature infant and babies with cardiac or respiratory compromise.

D. Technique

•

Prepare as for minor procedure (see Chapter 4). Ensure that neutral thermal environment is maintained. It is often necessary to transfer small babies to a radiant warmer for peripheral IV placement to avoid cold stress. If the infant has received a recent enteral feeding, consider delaying the procedure until before the next feeding or placing a nasal or oral gastric tube to empty the stomach in order to prevent aspiration.

- Use transillumination to visualize vessel if needed (see Chapter 12).
 - Select vessel for cannulation. The following is the suggested order of preference (see Fig. 26.2):
 - Back of handâ€"dorsal venous plexus
 - ∘ Forearmâ€"median antebrachial, accessory cephalic veins



FIG. 26.2. Simulated procedure showing IV needle held in dominant hand, while index finger and thumb of nondominant hand are used to anchor vein and stretch overlying skin.

- ∘ Scalp veinsâ€"supratrochlear, superficial temporal, posterior auricular
- o Footâ€"dorsal venous plexus
- Antecubital fossaâ€"basilic or cubital veins
- o Ankleâ€"small saphenous, great saphenous veins
- Cut hair with small scissors close to scalp if using scalp vessels. (Do not shave the area.)
- Warm limb with heel warmer, if necessary (rarely needed), for approximately 5 minutes.
- Apply tourniquet if anatomic site indicates.
 - Place as close to venipuncture site as possible.
 - Tighten until peripheral pulsation stops.
 - Release partially until arterial pulse is fully palpable.
- Prepare skin area with antiseptic. Allow to dry.

In the United States, povidone–iodine solution and isopropyl alcohol are the most commonly used skin disinfectant solutions. Povidone–iodine has been shown to have greater efficacy than isopropyl alcohol and, in addition, is less damaging to skin tissue. Povidone–iodine solution should be applied to the proposed insertion site and allowed to dry for at least 30 seconds. It then should be removed with sterile saline or sterile water. The importance of removing the povidone–iodine solution cannot be overstressed, as there have been reports of burns, elevated iodine levels, and hypothyroidism in premature infants caused by prolonged contact and further absorption (9).

- Attach syringe and T connector to needle/cannula, and test patency by forcing a small amount of saline through.
- Detach syringe and T connector.
- Select straight segment of vein or confluence of two tributaries.
- Grasp catheter between thumb and first finger. For winged angiocath, grasp plastic wings (Fig. 26.2).
- Anchor vein with index finger of free hand and stretch skin overlying it. This maneuver may also be used to produce distention of scalp veins.
- Hold needle parallel to vessel, in direction of blood flow.

- Introduce needle through skin, a few millimeters distal to point of entry into vessel (see Chapter 13).
- Introduce needle gently into vessel until blood appears in hub of needle or in cannula upon withdrawal of stylet.

When using a very small vessel or in an infant with poor peripheral circulation, blood may not appear immediately in tubing. Wait. If in doubt, inject a small amount of saline after releasing tourniquet.

- Remove stylet. Do not advance needle farther, because back wall of vessel may be pierced.
- Advance cannula as far as possible.

Injecting a small amount of blood or flush solution into the vein prior to advancing the cannula may assist cannulation (Fig. 26.3) (10).

- Remove tourniquet.
- Connect T connecter and syringe, and infuse small amount of saline gently to confirm intravascular position.
- Anchor needle or cannula as shown in Fig. 26.4



FIG. 26.3. Injecting a small amount of flush solution will distend wall of vein and facilitate cannulation. (Redrawn from Filston HC, Johnson DG. Percutaneous venous cannulation in neonates: a method for catheter insertion without œcutdown. Pediatrics 1971;48:896, with permission of American Academy of Pediatrics.)

- Attach IV tubing and secure to skin.
- If an armboard is necessary for securing site, place the affected extremity in an anatomically correct position before taping. Consider placing cotton or a 2 x 2-in gauze square beneath the hub of a T connector to prevent a pressure injury.



FIG. 26.4. Method for securing peripheral intravenous cannula with adhesive tape. A, B: Place an adhesive transparent tape over the cannula. C: Place tape 1 behind cannula as shown, with adhesive side up. D, E: Fold tape 1 anteriorly across the catheter–hub junction. F: Hold in place with tapes 2 and 3. The area of skin entry can be dressed with semipermeable sterile transparent dressing. Avoid obscuring with opaque dressing.

E. Complications (11,12 and 13)

- Hematoma the most common but not usually significant complication. Hematomas can often be managed with gentle manual pressure.
- Venospasm occurs rarely as a complication of venous access and usually resolves spontaneously
- Phlebitis (14,15)

Phlebitis remains the most common significant complication associated with the use of peripheral venous catheters. When phlebitis does occur, the risk of local catheter-related infection may be increased (15). Heparinized solutions (0.5 U/mL) have been shown to increase dwell time and reduce the frequency of complications such as phlebitis and erythema (16). The catheter material, catheter size, and tonicity of the infusate also influence the incidence of phlebitis. When peripheral lines are

used for parenteral nutrition, the coinfusion of a lipid solution with the hyperosmolar total parenteral nutrition solution prolongs the life of the vein (17,18).

- Infiltration of subcutaneous tissue with IV solution. (For management of this complication, see Chapter 27.) This is unfortunately a common complication of peripheral IV infusion. Extreme vigilance and avoidance of hyperosmolar IV solutions will help to reduce the incidence to the minimum possible.
 - Superficial blistering (Fig. 26.5)
 - Deep slough, which may require skin graft (Fig. 26.6)
 - Calcification of subcutaneous tissue due to infiltration of calcium-containing solution

Note that there may be some extravasation into adjacent tissues even though blood can be aspirated from the needle/cannula.

• Infection (19,20 and 21)

There is an increase in the incidence of both phlebitis and infection when a needle remains in place longer than 72 hours (7) and is heavily manipulated (21). An increase has also been reported with film-type dressings, but this remains controversial (1,2,3,4 and 5,15). Catheters made with Teflon or polyurethane appear to be associated with fewer infections in adults than catheters made with polyvinyl chloride or polyethylene (15). Polyurethane catheters appear to have an approximately 30% lower risk of phlebitis than Teflon catheters in adults (15). Batton et al. (14) failed to confirm a difference in the incidence of infection when 25-gauge needles were compared with 24-gauge Teflon cannulas. However, the Teflon cannulas remained functional three times as long as steel needles, with no apparent increase in complications.



FIG. 26.5. Result of infusion of lidocaine into subcutaneous tissues of lower limb.

- Embolization of clot with forcible flushing
- Hypernatremia, fluid overload, or heparinization of the infant due to improper flushing technique or solution; also electrolyte derangements from IV fluid infused at an incorrect rate



FIG. 26.6. Extensive deep skin slough that required grafting, caused by IV infiltration.



FIG. 26.7. A: Skin slough on scalp caused by inadvertent infusion into the frontal branch of the temporary artery. B: This is indicated by arrows.

- Accidental injection or infusion into artery with arteriospasm and possible tissue necrosis (Fig. 26.7)
- Burn from
 - Transilluminator (Fig. 26.8; also see Chapter 12)
 - Compress used to warm limb prior to procedure
 - ∘ Prolonged povidone–iodine or isopropyl alcohol application to very premature skin

- Air embolus
- Ischemia or gangrene of lower extremity, complicating infusion into saphenous vein; mechanism unclear (21)



FIG. 26.8. Burn from transilluminator used to locate vein in antecubital fossa.

Cutdown Placement of IV Catheter in the Great Saphenous Vein

Modern vascular access catheters and techniques have made the traditional cutdown largely obsolete. However, this means of IV access is occasionally required in emergency situations, particularly shock, when time is of the essence. The saphenous vein at the ankle is the safest, quickest site for the physician with limited surgical experience. Intraosseous vascular access is an alternative method of obtaining IV access in an emergency and is described in Chapter 47. The method described has the advantage of avoiding incision of the vessel prior to introduction of the catheter. This is an important advantage in the very small infant, in whom it is difficult to avoid excessive venotomy and transection of the vein. Even in the most experienced hands, the cutdown procedure may take 10 minutes and last no longer than percutaneous IV access. When other methods cannot be performed, venous cutdowns may provide the only alternative means of emergency venous access.

A. Indications

- To provide a route for peripheral IV therapy when the percutaneous method is not possible
- To provide a more stable and reliable IV line in situations where even brief cessation of therapy might compromise the infant
- To provide emergency IV therapy

B. Contraindications

- Risk/benefit should be weighed carefully in the presence of bleeding diathesis.
- Should not be used as routine procedure for starting IV when percutaneous method is technically difficult but not impossible.

C. Equipment Sterile

- Gown and gloves
- Cup with antiseptic solution (e.g., an iodophor)
- Sterile aperture drape
- 0.5% lidocaine HCl in 2-mL syringe
- Two 25-gauge venipuncture needles
- Two curved mosquito hemostats
- 22-gauge cannula with needle stylet or a short length of small-diameter (0.6- to 1.2-mm outer diameter) silicone rubber catheter. The small silicone catheters reduce irritation but can slow the process of placing the line.
- T connector for cannula
- Heparinized saline (for heparin lock).
- Half-strength normal saline in a 5-mL syringe
- Absorbable suture or 5.0 nylon suture on small, curved needle. It is preferable to close the wound with subcuticular absorbable sutures, whenever possible, to avoid inflammation and formation of suture tracts (6).
- Needle holder
- 11 scalpel blade and handle
- Semipermeable, sterile transparent dressing

Nonsterile

- Materials for restraint (see Chapter 3)
- Transilluminator (cover with sterile plastic glove to maintain sterile field; see Chapter 12)
- Roll of 0.5- to 1-in porous adhesive tape

D. Precautions

- Aspirate prior to injection of lidocaine to prevent inadvertent intravascular infusion.
- Take care not to make initial skin incision too deep, to avoid severing underlying vein.
- Avoid infusing extremely irritating or hypertonic solutions.

E. Technique

Anatomic considerations: The great saphenous vein is constant in its anatomic position, just anterior to the medial malleolus. It is the only structure of importance in this area. The cutdown procedure is facilitated by the fact that the vein lies on tough periosteum and has sufficient elasticity to allow withdrawal through a small incision without the danger of rupture.

- Restrain foot in equinovalgus position.
- Palpate medial malleolus, and locate point of incision 1 cm anterior and 1 cm superior to malleolus (Fig. 26.9).
- Scrub, put on mask, gown, and gloves, and prepare area of incision, as for major procedure (see Chapter 4).
- Drape area.
- Indicate line of incision by marking skin with sterile surgical pen prior to infiltration with local anesthetic.
- Infiltrate skin along line of incision with 0.5 to 1 mL of lidocaine, and then extend infiltration into subcutaneous tissue.
- Wait 5 minutes for anesthesia to take effect.



FIG. 26.9. Position of restraint for cutdown on the great saphenous vein at the ankle, indicating site of incision.



FIG. 26.10. Blades of curved hemostat are spread parallel to vein to dissect the subcutaneous connective tissue down to the periosteum.

- Make 1-cm transverse incision through skin, down to superficial subcutaneous fat. A vertical, rather than a transverse, incision is optional. The former has the advantage that it offers the opportunity to extend the incision cephalad, should the posterior wall of the vein be perforated on the initial attempt at cannulation. However, it has the disadvantage that it may be made too lateral or medial to the vein.
- Introduce curved hemostat into incision, with tip down. Spread blades of hemostat parallel to vein to dissect tissue down to periosteum. Continue this step until adequate visualization of vein is achieved (Fig. 26.10).



FIG. 26.11. A curved hemostat is used to "scoop†the vein into the incision.



FIG. 26.12. The hemostat has been carefully opened and the subcutaneous connective tissue spread, leaving the vein surface clean. A ligature is placed between the blades of the hemostat.

- Reintroduce curved hemostat into incision, with tip down, and pass down to periosteum. With a œscooping motion, through approximately 180 degrees, isolate vein and draw into incision (Fig. 26.11).
- Open hemostat carefully. Spread subcutaneous tissue, leaving vein surface clean.
- Place 5-0 silk suture loosely around vein and clamp at end of suture with hemostat to allow for distal control of vessel (Fig. 26.12). (Do not tie ligature.)
- Place ligature with clamp across extended index finger and inside palm of nondominant hand, retracting it in an upward and caudad direction (Fig. 26.13).



FIG. 26.13. Outward and caudad traction is exerted on the suture.



FIG. 26.14. Introducing the cannula into the vein.

- Introduce cannula/stylet into vein at a 45-degree angle, with bevel down. Once vein has been entered, angle cannula parallel to vein (Fig. 26.14).
- Advance cannula into vein while withdrawing inner needle stylet.



Α

FIG. 26.15. Cystogram in infant who had not urinated for more than 24 hours despite adequate intravenous fluids. A: The bladder appears normal, but there is a mass effect displacing the intestines in approximate area indicated by arrows. B: Radiographic contrast material, injected through a long catheter introduced into the femoral vein via the great saphenous vein, has extravasated into the abdominal cavity.

- Advance cannula up to hub, and infuse small volume of saline flush solution to confirm intravenous position.
- Remove traction suture and close skin incision with subcuticular absorbable sutures or one or two simple 5-0 nylon sutures.
- Attach cannula to infusion tubing and regulate IV.
- Secure cannula to skin as shown in Fig. 26.4.

F. Complications

- Same as for percutaneous method
- Inadvertent infusion of local anesthetic into artery or vein
- Severance of vein owing to excessively deep initial incision
- Infiltration of intravenous infusion into body cavity (Fig. 26.15)

This is a complication related to placement of very long catheters. When infusion of an extremely irritating or hypertonic solution is required, the catheter is preferably inserted into the central venous system (see Chapter 31).

• Varicose veins secondary to postinfusion phlebitis (22)

Conversion of Peripheral IV Line to a Heparin Lock

A. Technique

- Wash hands and put on gloves.
- Clean IV tubing and catheter connection with antiseptic solution.
- Stop IV infusion and remove IV tubing from hub of IV needle or cannula.
- Seal hub with a sterile plug or T-connector system (e.g., Argyle intermittent infusion plug [Consolidated Medical Equipment, Utica, NY, USA; Sherwood Medical Co., St. Louis, MO, USA]

or Burron spin-lock port extension set [Burron Medical, Bethlehem, PA, USA] that has been primed with the required quantity of heparinized saline). As an improvisation, a stopcock with two dead heads may be used. However, at least 3 mL of flush solution is necessary to flush all parts of a stopcock. This increases the margin for error, with possible fluid overload in very small premature infants.

- Clean plug with antiseptic, and inject 0.4 to 0.8 mL of heparinized saline solution through plug to flush blood from needle or cannula.
- Clean plug with antiseptic prior to every use.
- Refill heparin lock with heparinized flush solution after every IV infusion. (Flush routinely every 6 to 12 hours, depending on frequency of use.)

References

1. Wille JC, Blussae E, Vanovd Ablas A. A comparison of four film-type dressings by their antimicrobial effect on the flora of the skin. J Hosp Infect. 1989;14:153.

2. Vernon HJ, Lane AT, Wischerater LJ, et al. Semipermeable dressing and transepidermal water loss in premature infants. Pediatrics. 1990;86:357.

3. Maki DG, Ringer M. Evaluation of dressing regimens for prevention of infection with peripheral intravenous catheters. JAMA. 1987;258:3396.

4. Craven DE, Lichtenberg DA, Kunches LM, et al. A randomized study comparing a transparent polyurethane dressing to a dry gauze dressing for peripheral intravenous sites. Infect Control. 1985;6:361.
5. Holland KT, Harnby D, Peel B. A comparison of the in vivo antibacterial effects of "Op-Site,†â€œTegaderm†and "Ensure†dressings. J Hosp Infect. 1985;6:299.

6. Ganderer MW. Vascular access techniques and devices in the pediatric patient. Surg Clin North Am. 1992;72:1267.

7. Downing JW, Charles KK. Intravenous cannula fixing and dressingâ€" comparison between the use of transparent polyurethane dressing and conventional technique. South Afr Med J. 1987; 721:191.

8. Leibovici C. Daily change of an antiseptic dressing does not prevent infusion phlebitis: a controlled trial. Am J Infect Control. 1989;17:23.

9. Lund C, Kuller J, Lane A, et al. Neonatal skin care: the scientific basis for practice. JOGNN 1999;28:241. 10. Filston HC, Johnson DG. Percutaneous venous cannulation in neonates: a method for catheter insertion without "cut-down.†Pediatrics. 1971;48:896.

11. Johnson RV, Donn SM. Life span of intravenous cannulas in a neonatal intensive care unit. Am J Dis Child. 1988;142:968.

12. Duck S. Neonatal intravenous therapy. J Intravenous Nurs. 1997;20:121.

13. Wynsma L. Negative outcomes of intravascular therapy in infants and children. AACN Clin Issues. 1998;9:49.

14. Batton DG, Maisles JM, Appelbaum JM. Use of intravenous cannulas in preterm infants: a controlled study. Pediatrics. 1982;70:487.

15. Centers for Disease Control. Special communication. Guideline for prevention of intravascular device-related infections. Am J Infect Control. 1996;24:262.

16. Moclair A, Bates I. The efficacy of heparin in maintaining peripheral infusions in neonates. Eur J Paediatr. 1995;154:567.

17. Pineault M, Chessex P, Pledboeuf B, et al. Beneficial effect of coinfusing a lipid emulsion on venous patency. J Parenter Enter Nutr. 1989;13:637.

18. Phelps SJ, Lochrane EB. Effect of the continuous administration of fat emulsion on the infiltration rate of intravenous lines in infants receiving peripheral parenteral nutrition solutions. J Parenter Enter Nutr. 1989;13:628.

19. Lloyd-Still JD, Peter G, Lovejoy FH. Infected "scalp-vein†needles. JAMA. 1970;213:1496.
20. Lozon, MM. Pediatric vascular access and blood sampling techniques. In: Roberts JR, Hedges JR, eds. Clinical Procedures in Emergency Medicine. 4th ed. Philadelphia: Saunders; 2004:366–371.

Cronin WA, Germanson TP, Donowitz LG. Intravascular cannula colonization and related blood stream infection in critically ill neonates. Infect Control Hosp Epidemiol. 1990;11:301.
 Shuster S, Laks H. Varicose veins following ankle cut-downs. J Pediatr Surg. 1973;8:245.

27-Management of Extravasation Injuries

Jayashree Ramasethu

Extravasation or inadvertent infiltration of intravenously (IV)-administered solutions into subcutaneous tissue is a common adverse event in intensive care nurseries and may result in partial or complete skin loss, infection, and nerve and tendon damage, with the potential risk of cosmetic and functional impairment (1,2 and 3). Parenteral alimentation fluids, antibiotics such as nafcillin, calcium, potassium, and sodium bicarbonate solutions, and vasopressor agents are often implicated (1,4,5). Early identification and appropriate management are vital to minimize damage (6).

A. Assessment (Figs. 27.1; see also Figs. 26.5 and 26.6)

- Staging of extravasations is recommended for objective evaluation to determine the degree of intervention required, but detailed descriptions or digital photographs provide better documentation of the extent of the wound and the healing process (Table 27.1) (7,8).
- Fussiness, crying, or withdrawal of the limb when flushing the IV cannula are early warning signs, but these may be absent in an infant who is sedated or critically ill.
- Blistering and discoloration of skin often portend at least partial skin loss, but visible skin changes do not always indicate the severity of underlying injury, which may evolve over several days.

B. Management

The degree of intervention is determined by the stage of extravasation, the nature of the infiltrating solution, and the availability of specific antidotes. There is no consensus on management of Stage 3 or 4 lesions. In the absence of randomized controlled trials, some institutions have established management protocols to guide therapy, based on local experience, case series, and anecdotal evidence (1,7,8,9,10,11 and 12).

- In all cases:
 - Stop the IV infusion promptly.
 - Remove constricting bands that may act as tourniquets (e.g., armboard restraint).
 - Elevation of the limb may help to reduce edema.
 - The application of warm or cold packs is controversial. Warm packs may, by local vasodilation, help to reabsorb infiltrating solutions. However, warm moist packs have been reported to cause maceration of the skin (9).
- Stage 1 or 2 extravasation:
 - Remove IV cannula.
 - Consider antidote (see below).
- Stage 3 or 4 extravasation:
 - Leave the IV cannula in place and, using a 1-mL syringe, aspirate as much fluid as possible from the area.
 - Remove the cannula unless it is needed for administration of the antidote.
 - Consider use of hyaluronidase (4) or a specific antidote (see below).
 - Multiple-puncture technique (10): In infants who develop tense swelling of the site with blanching of the skin owing to infiltration of acidic or hyperosmolar solutions, multiple punctures of the edematous area using a blood-drawing stylet (and strict aseptic technique) has been used to allow free drainage of the infiltrating solution, decrease the swelling, and prevent necrosis. The area is then dressed with saline soaks to aid drainage.

- Saline flushout: A technique of saline flushing of the subcutaneous tissue has been advocated by some authors (2,6,11,12). After cleaning and infiltrating the area with 1% lignocaine, 500 to 1000 U of hyaluronidase is injected subcutaneously. Four small stab incisions are then made in the tissue plane with a scalpel blade at the periphery of the area. Saline is injected through a blunt cannula inserted subcutaneously through one of the puncture sites and flushed through the other puncture sites, massaging the fluid toward the incisions to facilitate removal of the extravasated material.
- Hyaluronidase: Dispersing agent effective in extravasations involving calcium, parenteral alimentation fluids, antibiotics, sodium bicarbonate, etc. Although standard reference manuals state that hyaluronidase is not recommended for treatment of vasopressor extravasation injury (13), there have been reports of successful treatment of such extravasations with a combination of hyaluronidase and saline irrigations, as described above (2,11).
 - Mechanism of action: Breakdown of hyaluronic acid, the ground substance or intercellular cement of tissues; minimizes tissue injury by enhancing dispersion and reabsorption of extravasated fluids

TABLE 27.1 Staging of Extravasation Injury (7,8)

StageCharacteristics

- Pain at siteâ€"crying when IV cannula is flushed IV cannula flushes with difficulty No redness or swelling
 Pain Redness and slight swelling at site Brisk capillary refill
 Pain Moderate swelling Blanching of area Skin cool to touch Brisk capillary refill below site Good pulse below site
- 4 Pain

Severe swelling around site Blanching of area Skin cool to touch Area of skin necrosis or blistering Prolonged capillary refill time (>4 s) Decreased or absent pulse



FIG. 27.1. A: Stage IV extravasation injury with blistering of skin. B: Same area 2 weeks later, with eschar formation.

- Most effective within 1 hour; may be used up to 12 hours
- Administration: Use 25- or 26-gauge needles to inject 1 mL (150 USP units/mL) as five separate 0.2-mL injections around the periphery of the extravasation site.
- Adverse effects: None reported in neonates, rare sensitivity reactions reported in adults
- Specific antidotes
 - Phentolamine (14,15)
 - Effective in treating extravasations of vasopressors such as dopamine and epinephrine, which cause tissue damage by intense vasoconstriction and ischemia
 - Effect should be seen almost immediately; most effective within 1 hour but may be used up to 12 hours. The biologic half-life of subcutaneous phentolamine is less than 20 minutes.
 - Mechanism of action: Competitive alpha-adrenergic blockade, leading to smooth muscle relaxation and hyperemia
 - Doses have not been established for newborn infants. The exact dose is dependent on the size of the lesion and the size of the infant.
 - Recommended doses range from 0.01 mg/kg per dose to 5 mL of 1-mg/mL solution (13,16).
 - Administration: 0.5 to 1 mg/mL of solution injected subcutaneously into infiltrated area, after removal of IV catheter
 - Precautions: Hypotension, tachycardia, and dysrythmias may occur; use with extreme caution in preterm infants; consider using repeated small doses.
 - Topical nitroglycerine (17,18)
 - Effective in treating injury due to extravasation of dopamine
 - Mechanism of action: Vascular smooth muscle relaxant
 - Application: 2% nitroglycerine ointment, 4 mm/kg body weight, applied over the affected area, may be repeated every 8 hours if perfusion has not improved (17)
 - Transderm–nitro patches have also been used (7).
 - Precautions: Absorption through the skin may lead to hypotension.
 - Terbutaline
 - Effective in treating peripheral ischemia resulting from extravasation of vasopressors in adults and older children; no neonatal cases reported (19)
 - Mechanism of action: Peripheral vasodilation by b₂-adrenoreceptor activation
 - Administration: Subcutaneous injection of terbutaline in concentration of 0.5 to 1 mg/mL; doses in adults vary from 0.5 to 1 mg.

- Wound management
- The goal of wound management in neonates who have partial- or full-thickness skin loss is to achieve primary or secondary healing while avoiding scarring, contractures, and operative intervention. Wound care regimens differ among experts and institutions (1,3,7,20).
 - Clean area with sterile saline.
 - Apply silver sulfadiazine cream to affected area and change dressing every 8 hours with gentle cleansing of wound and reapplication of cream. Sulphonamides increase the risk of kernicterus and therefore are contraindicated in infants in the first 30 days of life.
 - Amorphous hydrogels consisting of carboxymethylcellulose polymer, propylene glycol, and water have been shown to keep the wound moist and facilitate wound healing (20). They are available in the form of gels or sheets, which may be applied directly to the wound surface and held in place by a secondary dressing. The gel is easily removed with saline and is generally changed every 3 days. More frequent dressing changes may be required if there is excessive exudation.
 - Wet-to-dry saline dressings and povidone-iodine dressings may also be effective (9). The liberal use of povidone-iodine on open wounds is not recommended in very low-birthweight infants, because absorption of iodine from the skin may suppress thyroid function.
 - \circ $\;$ The role of antibacterial ointments is unclear.
 - Evaluate wound healing every day. Time to heal ranges from 7 days to 3 months.
 - If the scar involves a flexion crease, passive range-of-motion exercises with each diaper change may help to prevent contractures.
- Plastic surgical consultation
 - Recommended for all full-thickness and significant partial-thickness extravasation injuries
 - \circ Enzymatic or surgical dÃ[©] bridement or skin grafting may be required (3,21 and 23).

References

1. Wilkins CE, Emmerson AJB. Extravasation injuries in regional neonatal units. Arch Dis Child Fetal Neonatal Ed. 2004;89:

2. Casanova D, Bardot J, Magalon G. Emergency treatment of accidental infusion leakage in the newborn: report of 14 cases. Br J Plast Surg. 2001;54:396–399.

3. Friedman J. Plastic surgical problems in the neonatal intensive care unit. Clin Plast Surg. 1998;25:599–617.

4. Zenk KE, Dungy CI, Greene GR. Nafcillin extravasation injury. Use of hyaluronidase as an antidote. Am J Dis Child. 1981;135:1113–1114.

5. Chen JL, O'Shea M. Extravasation injury associated with low dose dopamine. Ann Pharmacother. 1998;32:545–548.

6. Gault DT. Extravasation injuries. Br J Plast Surg. 1993;46:91–96.

7. Flemmer L, Chan JSL. A pediatric protocol for management of extravasation injuries. Pediatr Nurs. 1993;19:355–358.

8. Montgomery LA, Hanrahan K, Kottman K. Guideline for IV infiltrations in pediatric patients. Pediatr Nurs. 1999;25:167–180.

9. Brown AS, Hoelzer DJ, Piercy SA. Skin necrosis from extravasation of intravenous fluids in children. Plast Reconstruct Surg. 1979;64:145–150.

10. Chandavasu O, Garrow D, Valda V, et al. A new method for the prevention of skin sloughs and necrosis secondary to intravenous infiltration. Am J Perinatol. 1986;3:4–6.

11. Harris PA, Bradley S, Moss ALH. Limiting the damage of iatrogenic extravasation injury in neonates. Plast Reconstruct Surg. 2001;107:893–894.

12. Davies J, Gault D, Buchdahl R. Preventing the scars of neonatal intensive care. Arch Dis Child. 1994;70: F50-F51.

13. Young TE, Magnum B. Neofax®: A Manual of Drugs Used in Neonatal Care. 18th ed. Raleigh, NC: Acorn; 2005:209.

14. Subhani M, Sridhar S, DeCristafaro JD. Phentolamine use in a neonate for the prevention of dermal necrosis caused by dopamine: a case report. J Perinatol. 2001;21:324-326.

15. Siwy BK, Sadove AM. Acute management of dopamine infiltration injury with regitine. Plast Reconstruct Surg. 1987;80:610–612.

16. Gomella TL, Cunningham MD, Eyal FG, Zenk KE. Neonatology: Management, Procedures, On-Call Problems, Diseases and Drugs. 5th ed. New York: Lange Medical Books/McGraw-Hill; 2004:629.

17. Wong AF, McCullough LM, Sola A. Treatment of peripheral tissue ischemia with topical nitroglycerine ointment in neonates. J Pediatr. 1992;121:980–983.

18. Denkler KA, Cohen BE. Reversal of dopamine extravasation injury with topical nitroglycerine ointment. Plast Reconstruct Surg. 1989;84:811–813.

19. Stier PA, Bogner MP, Webster K, et al. Use of subcutaneous terbutaline to reverse peripheral ischemia. Am J Emerg Med. 1999;17:91–94.

20. Cisler-Cahill L. A protocol for the use of amorphous hydrogel to support wound healing in neonatal patients: an adjunct to nursing care. Neonatal Network. 2006;25:267–273.

21. Falcone PA, Barrall DT, Jeyarajah DR, Grossman JAI. Nonoperative management of full thickness intravenous extravasation injuries in premature neonates using enzymatic debridement. Ann Plastic Surg. 1989;22:146–149.

22. Tiras U, Erdeve O, Karabulut AA, et al. Debridement via collagenase application in two neonates. Pediatr Dermatol. 2005; 22:472–475.

23. Schafer T, Kukies S, Stokes TH, et al. The prepuce as a donor site for reconstruction of an extravasation injury to the foot in a newborn. Ann Plast Surg. 2005;54:664–666.

28-Umbilical Artery Catheterization

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A. Indications

B.

Catheters should remain in place only as long as primary indications exist, with the exception of secondary indication A.3. Because of the risk of complications, catheters should usually not remain in place for more than 2 weeks.

Primary

- Frequent or continuous (see Chapter 9) measurement of lower aortic blood gases for oxygen tension (PO₂) or oxygen content (percent saturation)
- Continuous monitoring of arterial blood pressure
- Angiography
- Resuscitation (use of umbilical venous line may be first choice)

Secondary

• Infusion of maintenance glucose/electrolyte solutions or medications. If this line is to be used to provide intravenous nutrition, the same aseptic techniques must be used to prevent line-related sepsis as are used for any central line. See Chapter 31.

- Exchange transfusion
- To provide vital infusions (1) and a port for frequent blood sampling in the extremely lowbirthweight infant.

B. Contraindications

- Evidence of local vascular compromise in lower limbs or buttock areas
- Peritonitis
- Necrotizing enterocolitis (2)
- Omphalitis
- Omphalocele
- Acute abdomen etiology

C. Equipment

Several standardized graphs for premeasurement of catheter length to be inserted are available (Figs. 28.1 28.2 and 28.3).

Sterile

- Sterile gown and gloves
- Cup with antiseptic solution
- Surgical drape with central aperture (transparent drape recommended)
- Catheter
 - Single hole
 - Reduces surfaces for potential thrombus formation
 - Recorded pressure tracing will change when hole is occluded.
 - Made of flexible material that does not kink as it follows the curves of vessels
 - Relatively rigid walls with frequency characteristics suitable for accurate measurement of intravascular pressure
 - Small capacity (minimum volume of blood to be withdrawn to clear catheter prior to blood sampling)
 - Radio-opaque (the need to visualize the catheter position on x-ray film outweighs the theoretical risk of increased thrombogenicity related to a radio-opaque strip) (3)
 - Smooth, rounded tip (4), nonthrombogenic material (5)
 - 5-French (Fr) gauge for infants weighing >1,200 g
 - \circ 3.5-Fr gauge for infants weighing <1,200 g
- Scissors for cutting catheter
- Blunt needle adapter (if using catheter without hub)
 - No. 18 for 5-Fr catheter
 - No. 20 for 3.5-Fr catheter
- Three-way stopcock with Luer-Lok
- 10-mL syringe
- 0.25 to 0.5 N saline flush solution (saline with heparin, 1 to 2 U/mL)
- In very small premature infants, particularly in the first week of life, hypernatremia may result from receiving excess sodium in flush solutions. We recommend using 0.25 N rather than more concentrated saline solutions (6). Use of heparinized flush solution is common practice. Rajani and others (7,8) have shown that using a heparinized solution containing 1 U heparin/mL for flushing the umbilical arterial line prolonged catheter life by reducing the incidence of fibrin thrombus formation in the catheter lumen. Horgan et al. (9) found that the use of 1 U/mL heparin did not reduce the incidence of umbilical artery catheter (UAC)-related thrombi but did lower the incidence of their sequelae. Butt et al. (10) could demonstrate no significant benefit associated with increasing the rate of infusion from 1 to 2 mL/h (heparin 1 U/mL), and Bosque and Weaver (11) showed that continuous infusion of 1 U/mL heparin is more effective than intermittent infusion in maintaining

patency of the UAC. More recent data have indicated that heparin decreases the incidence of thrombotic complications (12), and a Cochrane Database Review found that the use of as little as 0.25 U/mL heparin in the infusate decreases the likelihood of line occlusion (13).

- Tape measure
- 20-cm narrow umbilical tie
- No. 11 scalpel blade and holder
- 4- x 4-in gauze sponges
- Two curved mosquito hemostats
- Toothed iris forceps
- Two curved, nontoothed iris forceps
- 2% lidocaine HCl without epinephrine
- 3-mL syringe and needle to draw up lidocaine
- Small needle holder
- 4-0 silk suture on small, curved needle
- Suture scissors



FIG. 28.1. Graph for determination of length of catheter to be inserted for appropriate low aortic or venous placement. Length of catheter is measured from umbilical ring. Length of umbilical stump must be added. The shoulder–umbilicus distance is the perpendicular distance between parallel horizontal lines at the level of the umbilicus and through the distal ends of the clavicles. (Adapted from

Dunn P. Localization of the umbilical catheter by postmortem measurement. Arch Dis Child. 1966; 41:69 , with permission)


FIG. 28.2. A: Graph for distance of catheter insertion from the umbilical ring for L3, L5, and aortic bifurcation. Large dots represent catheters positioned at L4. B: Graph for catheter insertion to level T8 using total body length. (From Rosenfeld W, Biagtan J, Schaeffer H, et al. Evaluation of graphs for insertion of umbilical artery catheters below the diaphragm. J Pediatr. 1981;98:628, with permission.)

Nonsterile

- Cap and mask
- Wooden tongue depressor

D. Precautions

- Avoid use of feeding tubes as catheter (associated with higher incidence of thrombosis) (14).
- Fold drapes so as not to obscure infant's face and upper chest.
- Take time and care to dilate lumen artery before attempting to insert catheter.
- Catheter should not be forced past an obstruction.
- Never advance catheter once placed and secured.
- Loosen umbilical tie slightly upon completion of procedure and obtain radiographic confirmation of position.
- Avoid covering the umbilicus with dressing. Dressing may delay recognition of bleeding or catheter displacement.
- Always obtain radiographic or ultrasound (15,16) confirmation of catheter position. If doubt remains, obtain a lateral radiographic study (17,18).
- Be certain that catheter is secure, and examine frequently when infant is placed in prone position, because hemorrhage may go unrecognized.
- Take care not to allow air to enter the catheter. Always have catheter fluid filled and attached to closed stopcock prior to insertion. Check for air bubbles in catheter before flushing or starting infusion.
- When removing catheter, cut suture at skin, not on catheter, to avoid catheter transection.



FIG. 28.3. Estimates of insertion length of umbilical catheters (umbilical artery catheter tip inserted between T6 and T10; umbilical vein catheter tip inserted above diaphragm in inferior vena cava near right atrium) based on birthweight (BW) (with 95% confidence intervals). Modified estimating equations utilizing BW are as follows: umbilical artery length = 2.5 BW + 9.7 (top) and umbilical vein length = 1.5 BW 5.6 (bottom), where BW is measured in kilograms and lengths are measured in centimeters. (From Shukla H, Ferrara A. Rapid estimation of insertional length of umbilical catheters in newborns. Am J Dis Child. 1986;140:787, with permission.)

E. Technique (See also Umbilical Catheterization on the Procedures DVD)

Anatomic note: The umbilical arteries are the direct continuation of the internal iliac arteries. Their diameters at their origins are 2 to 3 mm. As they approach the umbilicus, their lumina become small and the walls thicken significantly. In a full-term infant, each artery is approximately 7.0 cm long (Fig. 28.4). A catheter introduced into the umbilical artery will usually pass into the aorta from the internal iliac artery. Occasionally, it will pass into the femoral artery via the external iliac artery or into one of the gluteal arteries. The latter two sites are unsuitable for sampling, pressure measurement, or infusion.

• Choose either of two positions (Fig. 28.5).

High position is associated with fewer episodes of blanching and cyanosis of the lower extremities (19). High catheters were found to have decreased incidence of clinical vascular complications with a relative risk of 0.53 (95% confidence interval, 0.44 to 0.63) with no statistically significant increase in any adverse sequelae, including the incidence of hypertension, intraventricular hemorrhage, hematuria, necrotizing enterocolitis, or death (20).

- Low position (21,22 and 23): Level of lumbar vertebrae 3 to 4 (Fig. 28.6)
 - Catheter tip is below major aortic branches such as renal mesenteric arteries.
 - In most newborns, this position coincides with the aortic bifurcation at the upper end of the fourth vertebra.
- High position (14,19,24): Level of thoracic vertebrae 6 to 9 (Fig. 28.7); catheter tip above origin of celiac axis
- Make external measurements as necessary to estimate length of catheter to be inserted (see Figs. 28.1,28.2 and 28.3) (25,26,27,28,29,30 and 31).
- Prepare as for major procedure (see Chapter 4).



FIG. 28.4. Anatomic relations of the umbilical arteries, showing relationships with major arteries supplying buttocks and lower limb.



FIG. 28.5. The aorta and branches.

- Attach stopcock to hub of catheter and fill system with flush solution. Turn stopcock "off†to catheter. If catheter does not have a hub, cut flared end of catheter with scissors and insert blunt needle adapter of appropriate size. This will reduce catheter dead space.
- Place sterile gauze around umbilical stump and elevate out of sterile field, or have ungloved assistant grasp cord by cord clamp or forceps and pull cord vertically out of sterile field.
- Prepare cord and surrounding skin with antiseptic solution to radius of approximately 5 cm. The use of chlorhexidine in infants <2 months of age is not suggested (32).
- Drape area surrounding cord.
- Place umbilical tie around umbilicus and tie loosely with a single knot.

- Tighten only enough to prevent bleeding and place, if possible, around Wharton jelly rather than skin.
- \circ $\;$ It may be necessary to loosen the tie when inserting the catheter.



FIG. 28.6. Anteroposterior (A) and lateral (B) radiographs showing optimal low position of an umbilical artery catheter. Catheter tip is at the level of the superior margin of the fourth lumbar vertebral body, which in newborns usually corresponds to the aortic bifurcation.

- Cut cord horizontally with scalpel (Fig. 28.8).
 - Approximately 1 to 1.5 cm from skin
 - Avoid tangential slice.

Bloom et al. (33) have described an alternative approach to the artery with lateral arteriotomy. To perform this method, 3 to 4 cm of cord must be preserved because the cord must be rolled over a Kelly clamp 180 degrees (33,34).

- Clamp across end of cord with a mosquito hemostat in the nondominant hand and pull firmly toward the infant's head.
- Roll cord 180 degrees over hemostat toward abdominal wall.
- Identify arteries in superior right and left lateral aspects of cord.
- Approximately 1 cm from abdominal wall, incise Wharton jelly down to arterial wall, using no. 11 scalpel blade.
- Incise artery through half of circumference. If necessary, dilate lumen with iris forceps.
- Insert catheter into lumen of artery, directed in a caudad direction, for predetermined distance.
- Control bleeding by gentle tension on umbilical tape.
- Blot surface of cord stump with gauze swab. Avoid rubbing, because this damages tissue and obscures anatomy.



FIG. 28.7. Umbilical arterial catheter in satisfactory high position at the level of the ninth thoracic vertebral body on anteroposterior (A) and lateral (B) projections.

- Identify cord vessels (Fig. 28.9).
 - Vein is easiest to identify as large, thin-walled, sometimes gaping vessel. It is most frequently situated at the 12 o'clock position at the base of the umbilical stump.
 - Arteries are smaller, thick-walled, and white and may protrude slightly from cut surface.
 - Omphalomesenteric duct is rarely present.



FIG. 28.8. Traction is being placed on cord in direction of the arrow. Operator is about to make a horizontal cut across cord.



FIG. 28.9. The vessels of the umbilical cord. Thin-walled umbilical vein at 12 o'clock position is indicated by a white arrow. One of the two umbilical arteries is to the right and directly below the vein.

- Grasp cord stump, using toothed forceps, at point close to (but not on) artery to be catheterized. If available, it may be helpful to have an assistant scrub and assist.
 - Apply two curved mosquito hemostats to Wharton jelly on opposite sides of cord, away from vessel to be cannulated.
 - Apply traction to stabilize cord stump.



FIG. 28.10. An iris forceps is pointed into the umbilical artery in order to dilate the lumen of the artery.



FIG. 28.11. A: Inserting the catheter into the artery between the prongs of dilating forceps. Note that the umbilical tape has been tied around the skin of the umbilicus; this should be loosened once the catheter is secured in place. B: Close-up photo of the umbilical stump with the arterial catheter in place.

- Introduce one of the points of the curved iris forceps into the lumen of the artery and probe gently to a depth of 0.5 cm.
- Remove forceps and bring points together before introducing them once more into the lumen.
- Probe gently to a depth of 1 cm (up to the curved "shoulder†of the forceps), keeping the points together.
- Allow the points to spring apart, and maintain forceps in this position for 15 to 30 seconds to dilate vessel (Fig. 28.10). Time spent in ensuring dilatation prior to catheter insertion increases the likelihood of success.
- Release cord and set aside toothed forceps, while keeping curved forceps within artery.
- Grasp catheter 1 cm from tip, between free thumb and forefinger or with curved iris forceps.
- Insert catheter into lumen of artery, between prongs of dilating forceps (Fig. 28.11).
- Remove curved forceps, having passed catheter approximately 2 cm into vessel with a firm, steady motion. Grasp cord again with toothed tissue forceps and pull gently toward head of infant. This mild traction will facilitate passage of catheter at an angle between the cord and the abdominal wall.
- After passing the catheter approximately 5 cm, aspirate to verify intraluminal position. Clear blood by injecting 0.5 mL of flush solution. The catheter may now be used to measure blood gases.
- Take appropriate action if insertion is complicated (Fig. 28.12).
 - Resistance before tip reaches abdominal wall (<3 cm from surface of abdominal stump)
 - Loosen umbilical tape.
 - Redilate artery.
 - Popping sensation rather than relaxation



FIG. 28.12. Some reasons for failure of umbilical artery catheterization. A: Sagittal midline section to show normal anatomy of umbilical artery. B: Catheter has perforated the umbilical artery within the anulus umbilicalis and is dissecting perivascularly and external to peritoneum. C: Catheter has ruptured through the tunica intima (t.i.) and dissected into subintimal space. D: Catheter invaginating the tunica intima after stripping it from a more distal point. (Adapted from Clark JM, Jung AL. Umbilical artery catheterization by a cut down procedure. Pediatrics (Neonatol Suppl). 1977;59:1036, with permission of American Academy of Pediatrics.)

- Catheter may have exited lumen and created a false channel.
- Remove and use a second artery.
- If unsuccessful, draw 0.5 mL of lidocaine from vial. Reinsert tip of catheter approximately 2 cm into UAC and drip lidocaine into vessel. Apply constant gentle pressure until vessel dilates.
- o Backflow of blood, particularly around vessel
 - Tighten umbilical tape.
 - Catheter may be in false channel, with extravascular bleeding.
- Resistance is encountered at anterior abdominal wall or sharp turn in vessel as it angles around bladder toward internal iliac artery (approximately 6 to 8 cm from surface of umbilical stump in 2- to 4-kg neonate).
 - Apply gentle but steady pressure for 30 to 60 seconds.

- Position infant on side with same side elevated as artery being catheterized. Flex hip.
 Instill lidocaine as for 23.b (3). Do not force catheter.
- Easy insertion, but no blood return
 - Catheter is outside vessel in false channel.
 - Remove and observe infant carefully for evidence of complication.
- Place marker tape on catheter with base of tape flush with surface of cord, so that displacement of the catheter may be readily recognized.
- Remove umbilical tape and place purse-string suture around base of the cord (not through skin or vessels). Three bites into cord (with needle facing away from catheter) are sufficient to include all three vessels within the suture.

If desired, form marker tape into bilateral wings, and sew the tails of the purse-string suture through the wings to anchor the catheter in a symmetrical fashion. This is a useful method in very small premature infants because it avoids sticking tape to the abdominal wall (35). Or remove needle and wrap ends of suture in opposite direction around catheter for about 3 cm and tie, taking care not to kink catheter.

- Secure catheter temporarily by looping over upper abdomen and taping.
- Obtain radiograph or ultrasound (see D.8) to check catheter position.
 - \circ Catheter tip above T6 or between T11 and L2
 - Measure distance between actual and appropriate position on radiograph.
 - Withdraw equal length of catheter.
 - Repeat radiographic study.
 - Note procedure in chart.
 - Catheter tip below L5
 - Remove catheter.
 - Never advance catheter once in situ, because this will introduce a length of contaminated catheter into the vessel.
- If desired, secure catheter with tape bridge (Fig. 36.14).
- Continue routine cord care with triple dye or other agent of choice.
 - Stabilize catheter, stopcock, and syringe, using tongue depressor (optional).
 - Reduces risk of air embolus if syringe is maintained in vertical position
 - o Prevents accidental disconnection of catheter system

F. Alternative Technique: Umbilical Artery Cutdown

This method is usually successful even after failed insertion through the umbilical stump, as there is less tendency for false tracts. The most frequent reason for failed umbilical artery cutdown is mistaking the urachus for a vessel. Because of the time and risks associated with the cutdown procedure, standard insertion should be attempted first.

Indications

• Failed umbilical artery catheterization through conventional technique described earlier in this chapter

Contraindications

- Same as for umbilical artery catheterization by conventional technique
- Bleeding diathesis

Equipment

• Same as for umbilical artery catheterization by conventional technique.

- 1% lidocaine HCl without epinephrine in 3-mL syringe with 25- to 27-gauge needle
- No. 15 surgical blade and holder
- Curved delicate dressing forceps, two pairs (1/4 or 1/2 curved)
- Tissue forceps
- Self-retaining retractor (such as eyelid retractor)
- Absorbable suture, plain
- Absorbable suture on small cutting needle
- Nonabsorbable suture on a small, curved needle
- Needle holder
- Suture scissors
- Skin-closure tapes

Precautions

- Same as described earlier for conventional technique.
- If possible, leave catheter from previously attempted standard procedure in place to aid in vessel identification.
- Ensure that abdominal incision is on abdominal wall and not too close to umbilical stump.
- Identify landmarks carefully to avoid cutting or catheterizing urachus.
- When incising mesenchymal sheath, take care to avoid transecting vessel.
- Secure the catheter with an internal ligature that is just tight enough to prevent accidental removal but loose enough for elective removal or reinsertion in case the catheter becomes occluded by thrombus or precipitate.

Technique (36,37)

- See Fig. 28.13.
- Insert an oral gastric tube to keep the bowel as decompressed as possible.
- Prepare infant and drape as for umbilical artery catheterization (see earlier in chapter).
- If catheter has been left in place after previous attempt, include vessel and catheter in the preparation, leaving the catheter accessible for removal.
- Anesthetize area of skin immediately below umbilicus, at umbilical stump–abdominal wall junction, with 0.5 mL of lidocaine.
- Prepare umbilical artery catheter as for standard procedure, leaving catheter filled with flush solution. Estimate length for insertion based on patient size. Subtract 1 to 2 cm from that recommended for standard insertion, as cutdown catheter will enter vessel farther along course.
- Make a smile-shaped incision from 4 to 8 o'clock through the skin of the abdominal wall at the junction with the umbilical stump.
- Place self-retaining retractor to maintain exposure.
- Using blunt dissection through the subcutaneous tissue with mosquito forceps, identify the fascia overlying the urachus and umbilical vessels.

The mesenchymal sheath is composed of three layers of fascia and is from 1 to 3 mm thick. Although it is barely perceptible in extremely premature infants, in term infants it may be thick enough to require making an incision through the sheath prior to blunt dissection.

- While elevating the fascia with two forceps, make a small incision between their tips. Enlarge incision with scissors to the same size as skin incision. In very immature infants, simple dissection should suffice.
- With curved mosquito forceps, dissect in the midline and identify the urachus (Fig. 28.13).



FIG. 28.13. Subumbilical cutdown. Anatomic view through incision. (Redrawn from Sherman NJ. Umbilical artery cutdown. J Pediatr Surg. 1977;12:723, with permission.)

- The urachus is a white, glistening, cordlike structure in the midline. Its position may be confirmed by traction cephalad, pulling the dome of the bladder into view. The umbilical arteries lie posterolaterally on either side but not touching the urachus.
- Identify the umbilical arteries lying to either side of the urachus.

The vessels with their surrounding tissues appear larger than expected. When elevated, there will be no caudal bulge, distinguishing them from the urachus. If a previously attempted catheter was left in place, palpation of the area allows more ready identification of the vessel. Previously unsuccessful attempts, with failure to pass more than a few centimeters, are usually associated with perivascular hematoma formation from unrecognized perforation and dissection through a false tract. Visualization of a hematoma helps distinguish the vessel from the urachus.

- Try to avoid entering the peritoneum. In infants with very little subcutaneous tissue, it may be impossible to avoid penetrating the peritoneum. Should this occur, replace any bowel that may protrude and carefully close the peritoneum with absorbable suture, taking extreme care not to include any bowel within the suture. Start antibiotics for peritonitis prophylaxis.
- Insert the tip of the mosquito forceps under the vessel and pull a doubled strand of plain absorbable suture under the vessel. Position sutures 1 cm apart.
- While elevating the sutures and with suture scissors directed cephalad, make a V-shaped incision through three fourths of the diameter of the vessel. Take care not to transect the vessel, but cut cleanly into the lumen.

If the artery is accidentally transected and if the catheter insertion is unsuccessful, tie off the caudal end of the artery to prevent hemorrhage.

- Use curved tissue forceps or a catheter introducer to dilate the artery.
- Pass the catheter through the opening for the predetermined distance, checking for blood return after a few centimeters. The catheter should advance without resistance.

- When the catheter is properly positioned, have an assistant check the perfusion in the lower extremities. If that is satisfactory, secure the catheter by tying the lower ligature firmly around the catheter.
- Using absorbable suture, close the fascia and approximate the subcutaneous tissues.

Hashimoto et al. (38) proposed an alternative technique that allows for catheter reinsertion in case of catheter thrombosis or occlusion. They use loose ligation around the artery once the catheter is in proper position. They then fix the artery by using the same sutures that close the fascia, thus creating an arteriocutaneous fistula, making it easy to find the insertion site and use it for reinsertion.

- Close the skin with nonabsorbable suture or with skin-closure tape after cleaning the area.
- The catheter may be further secured with a tape bridge (Fig. 36.14).

Removal of catheter

- Remove any tape and withdraw catheter slowly as described earlier in this chapter.
- If the internal ligature around a catheter is too tight to allow removal with reasonable traction, it may be necessary to dissect and cut the ligature after sterile skin preparation.
- Apply pressure for hemostasis.
- Approximate wound edges with skin-closure tape.

Complications

- Catheterization of urachus (39)
- Vesicoumbilical fistula (39)
- Transection of urachus with urinary ascites (40)
- Perforation (41,42 and 43) or rupture (44) of urinary bladderâ€" although Nagarajan (45) has suggested that the risk of bladder injury is minimal if bladder is emptied prior to procedure.
- Transection of umbilical artery with hemorrhage
- Incision of peritoneum (with possible evisceration)
- Bleeding from incision

G. Care of Dwelling Catheter

For setup and maintenance of arterial pressure transducer, see Chapter 8.

- Keep catheter free of blood to prevent clot formation.
 - Flush catheter with 0.5 mL of flush solution, slowly over at least 5 seconds, each time a blood sample is drawn.
 - Between samples, infuse intravenous solution continuously through catheter to prevent retrograde flow.
 - Note amounts of blood removed and intravenous fluid/flush solution infused, and add to fluid balance record.
- Watch for indications of clot formation.
 - Decrease in amplitude of pulse pressure on blood pressure tracing
 - Difficulty withdrawing blood samples
- Take appropriate action if clot forms.
 - Do not attempt to flush clot forcibly.

Remove catheter. Replace only if critical line.

If further assistance is needed, the International Children's Thrombophilia Network is available to answer questions at 1-800-NO-CLOTS.

• Avoid enteral feedings with catheter in situ if possible. Increased risk of mesenteric thromboembolism has been suggested (46).

H. Obtaining Blood Samples from Catheter

(With emphasis on aseptic technique and minimizing stress to the vessel) Equipment

- Gloves
- Alcohol swabs
- Rubber-tipped clamps or disposable intravenous tubing clamps
- Syringe of 0.6 mL of flushing solution
- Syringe for cleaning line
- Syringe for blood sample
- Ice, if necessary for sample preservation
- Appropriate requisition slips and labels

Technique

- Wash hands and put on sterile gloves.
- Form sterile field.
- Clean the connection site of the stopcock/catheter using an alcohol swab.
- Clamp the umbilical catheter.
- Connect the 3-mL syringe, release the clamp, and slowly draw back 2 to 3 mL of fluid over 1 minute to clear the line. Reclamp the catheter. Remove syringe and place on sterile field. Data published by Davies et al. (47) indicate that accurate measurements of electrolytes can be obtained after withdrawal of a minimum of 1.6 mL of blood. However, if blood glucose values are desired, a minimum of 3 mL from a 3.5-Fr and 4 mL from a 5-Fr catheter must be withdrawn.
- Attach sampling syringe. Release clamp and draw back specimen desired. Reclamp the catheter.
- Reattach the syringe containing the fluid and blood cleared from the line.
 - Clear the connection of air.
 - Slowly replace the fluid and blood cleared from the line and remove the syringe.



FIG. 28.14. Various umbilical artery catheter (UAC) malpositions. A: Unacceptable position at L2 because of the proximity of the renal arteries. B: UAC in left brachycephalic artery. C: UAC in right brachycephalic artery. D: UAC in pelvic artery.

- Attach the syringe of flushing solution to the stopcock, clear air from the connection, and slowly flush the line.
- Clean the stopcock connection with alcohol.
- Record on infant's daily record sheet all blood removed and volume of flush used.

A study was carried out that looked at cerebral oxygenation and blood sampling from UAC in high position preterm infants (median gestational age 30 weeks). Although the clinical significance is unclear, the study showed that blood sampling of 2.3 mL (including flush volumes) through the UAC within 20 seconds resulted in a significantly decreased cerebral oxygenated hemoglobin and tissue oxygenation index. It also caused an increase in deoxygenated hemoglobin. This was not seen when the sampling time was extended to 40 seconds (48).

I. Removal of Umbilical Artery Catheter



FIG. 28.15. Anteroposterior (A) and lateral (B) radiographs demonstrating passage of an umbilical artery catheter into the pulmonary artery via a patent ductus arteriosus.

Indications

- No further clinical indication
- Need for less frequent direct PO₂ measurements
- Sufficient stabilization of blood pressure to allow intermittent monitoring
- Hypertension
- Hematuria not due to other recognizable cause
- Catheter-related sepsis and/or infections with Staphylococcus aureus, gram-negative bacilli, or Candida mandate removal of the catheter (49)
- Catheter-related vascular compromise
- Onset of platelet consumption coagulopathy
- Peritonitis
- Necrotizing enterocolitis
- Omphalitis

Technique

• Leave umbilical tie loose around cord stump as precaution against excessive bleeding.

Reinsertion of purse-string suture through dried Wharton jelly is preferable:

- Umbilical tape must be tied on skin rather than Wharton jelly.
- Catheter has been in situ for longer than 48 hours, because artery may have lost ability to spasm.
- Withdraw catheter slowly and evenly, until approximately 5 cm remains in vessel, tightening pursestring suture or umbilical tie.

- Discontinue infusion.
- Pull remainder of catheter out of the vessel at rate of 1 cm/min (to allow vasospasm). If there is bleeding, apply lateral pressure to the cord by compressing between thumb and first finger.

J. Complications (50,51,52,53,54 and 55)

Catheterization of the umbilical artery is probably always associated with some degree of reversible damage to the arterial intima (56,57).

- Malpositioned catheter (Figs. 28.14, 28.15 and 28.16) (46,58,59)
 - Vessel perforation (58,60,61)
 - Refractory hypoglycemia with catheter tip opposite celiac axis (62,63 and 64)
 - Peritoneal perforation (65)
 - False aneurysm (66,67,68,69,70 and 71)
 - Movement of catheter tip position because of changes in abdominal circumference
 - Sciatic nerve palsy (72,73)
 - Misdirection of catheter into internal or external iliac artery (see Figs. 28.14D and 28.17) (50).

Schreiber et al. (56) have described a double-arterial catheter technique to correct this problem.

- Vascular accident
 - Thrombosis (Fig. 28.18) (21,74,75,76,77 and 78)
 - Embolism/infarction (Fig. 28.17) (18,35,79,80,81 and 82) seen days or weeks after line insertion (83)
 - Vasospasm (18,83,84,85,86,87 and 88) is seen within minutes to a few hours after insertion.
 - Loss of extremity (Fig. 28.19) (86)
 - Hypertension (Fig. 28.20) (19,89,90,91,92,93,94 and 95)
 - Paraplegia (96,97,98,99,100 and 101)
 - Congestive heart failure (aortic thrombosis) (100)
 - Air embolism (Fig. 28.21)



FIG. 28.16. Effect of abdominal mass stimulating catheter misplacement. Anteroposterior (A) and lateral (B) films show remarkable displacement of an umbilical artery catheter by a giant hematocolpos in a 1-day-old infant.



A

FIG. 28.17. Vascular compromise in the left buttock and loin owing to a complication of umbilical artery catheter displaced into the internal iliac artery. For vascular anatomy, see Fig. 28.4.



FIG. 28.18. Arrows indicate mural thrombus in the abdominal aorta, which was associated with an umbilical arterial line. Upon further dissection of this autopsy specimen, the left renal artery was found to be occluded by thrombus. The left kidney is showing a degree of atrophy. Both kidneys showed scattered infarction.

- Equipment-related
 - Breaks in catheter and transection of catheter (58,85,97,102,103,104 and 105)
 - Plasticizer in tissues (60,104)
 - Electrical hazard (77)
 - Improper grounding of electronic equipment
 - Conduction of current through fluid-filled catheter
 - Intravascular knot in catheter (105)
- Other
 - Hemorrhage (including that related to catheter loss or disconnection and overheparinization) (50,79,85, 106,107 and 108)
 - o Infection (25,80,85,109,110,111,112,113,114,115,116,117 and 118)
 - Necrotizing enterocolitis (46,86)
 - Intestinal necrosis or perforation (119)
 - Vascular accident
 - Infusion of hypertonic solution (120)
 - Transection of omphalocele (Fig. 28.22) (121)
 - Herniation of appendix through umbilical ring (122)
 - Cotton fiber embolus (123)
 - Wharton jelly embolus (124)
 - Hypernatremia
 - True (6)
 - Factitious (104)
 - Factitious hyperkalemia (104)
 - Bladder injury (ascites) (125,126 and 127)
 - Curving back of the catheter on itself as a result of it catching in the intima (128)

- m Pseudo-coarctation of the aorta (129,130)
- Pseudo-mass in left atrium (131)
- Displacement by thoracoabdominal abnormality (132)



FIG. 28.19. Autoamputation of forefoot, owing to vascular complication of umbilical artery catheter.



FIG. 28.20. Generalized mottling of skin in infant with severe hypertension secondary to umbilical artery catheter-associated thrombus in renal artery.



FIG. 28.21. Anteroposterior roentgenogram demonstrating air embolism from an umbilical artery catheter in the left subclavian artery (upper arrow) and the femoral arteries (lower arrows).



FIG. 28.22. Small omphalocele. This gut-containing hernia was transected during placement of umbilical artery catheter.

References

1. Kanarek SK, Kuznicki MB, Blair RC. Infusion of total parenteral nutrition via the umbilical artery. J Parenter Enter Nutr. 1991;15:71.

2. Rand T, Weninger M, Kohlhauser C, et al. Effects of umbilical arterial catheterization on mesenteric hemodynamics. Pediatr Radiol. 1996;26:435.

3. Clawson CC, Boros SJ. Surface morphology of polyvinyl chloride and silicone elastomere umbilical artery catheters by scanning electron microscopy. Pediatrics. 1978;62:702.

4. Hecker JF. Thrombogenicity of tips of umbilical catheters. Pediatrics. 1981;67:467.

5. Boros SJ, Thompson TR, Reynolds JW, et al. Reduced thrombus formation with silicone elastomere (Silastic) umbilical artery catheters. Pediatrics. 1975;56:981.

6. Hayden WR. Hypernatremia due to heparinized saline infusion through a radial artery catheter in a very low birth weight infant. J Pediatr. 1978;92:1025.

7. Rajani K, Goetzman BW, Wennberg RP, et al. Effects of heparinization of fluids infused through an umbilical artery catheter on catheter patency and frequency of complications. Pediatrics. 1979;63:552.

8. Merenstein GB. Heparinized catheters and coagulation studies. J Pediatr. 1971;79:117.

9. Horgan MJ, Bartoletti A, Polansky S, et al. Effect of heparin infusates in umbilical arterial catheters on frequency of thrombotic complications. J Pediatr. 1987;111:774.

10. Butt W, Shann F, McDonnell G, et al. Effect of heparin concentration and infusion rate on the patency of arterial catheters. Crit Care Med. 1987;15:230.

11. Bosque E, Weaver L. Continuous versus intermittent heparin infusion of umbilical artery catheters in the newborn infant. J Pediatr. 1986;108:141.

12. Hentschel R, Weislock U, Von Lengerk C, et al. Coagulation-associated complications of indwelling arterial and central venous catheters during heparin prophylaxis: a prospective study. Eur J Pediatr. 1999;158(suppl 3):S126.

13. Barrington KJ, Umbilical artery catheters in the newborn: effects of heparin. Cochrane Database Syst Rev. 2000;(2): CD000507.

14. Westrom G, Finstrom O, Stenport G. Umbilical artery catheterization in newborns: thrombosis in relation to catheter tip and position. Acta Paediatr Scand. 1979;68:575.

15. Oppenheimer DA, Carroll BA, Garth KE, et al. Sonographic location of neonatal umbilical catheters. AJR. 1982; 138:1025.

16. Madan RJ, Deshpandes A. Reappraisal of ultrasound imaging of neonatal intravascular catheters. Arch Dis Child Fetal Neonat Ed. 1996;75:F622.

17. Baker DH, Berdon WE, James LS. Proper localization of umbilical arterial and venous catheters by lateral roentgenograms. Pediatrics. 1969;43:34.

18. Weber AL, Deluce S, Shannon DL. Normal and abnormal position of umbilical artery and venous catheter on the roentgenogram and review of complications. AJR. 1974;20:361.

19. Mokrohisky ST, Levine RL, Blumhagen RD, et al. Low positioning of umbilical artery catheters increases associated complications in newborn infants. N Engl J Med. 1978;229:561.

20. Barrington KJ. Umbilical artery catheters in the newborn: effects of position of the catheter tip. Cochrane Database Syst Rev. 2000;(2):CD000505.

21. Goetzman BW, Stadalnick RC, Bogen HG, et al. Thrombotic complications of umbilical artery catheters: a clinical and radiographic study. Pediatrics. 1975;56:374.

22. Neal WA, Reynolds JW, Jarvis CW, et al. Umbilical artery catheterization: demonstration of arterial thrombosis by aortography. Pediatrics. 1972;50:6.

23. Tooley WH. What is the risk of an umbilical artery catheter? Pediatrics. 1972;50:1.

24. Avery ME, Fletcher BD. The Lung and Its Disorders in the Newborn Infant. Philadelphia: WB Saunders; 1974.

25. Adam RD, Edwards LD, Becker CC, et al. Semi-quantitative cultures and routine tip cultures on umbilical catheters. J Pediatr. 1982;100:123.

26. Dunn P. Localization of the umbilical catheter by post-mortem measurement. Arch Dis Child. 1966;41:69.

27. Rosenfeld W, Biagtan J, Schaeffer H, et al. A new graph for insertion of umbilical artery catheters. J Pediatr. 1980;96:735.

28. Rosenfeld W, Estrada R, Jhaveri R, et al. Evaluation of graphs for insertion of umbilical artery catheters below the diaphragm. J Pediatr. 1981;98:627.

29. Shukla H, Ferrara A. Rapid estimation of insertional length of umbilical catheters in newborns. Am J Dis Child. 1986; 140:786.

30. Weaver R, Ahlgren E. Umbilical artery catheterization in neonates. Am J Dis Child. 1971;122:499.

31. Rubin BK, McRobert E, O'Neil MB. An alternate technique to determine umbilical arterial catheter length. Clin Pediatr. 1986;25:407.

32. Latini G. Potential hazards of exposure to di-(2-ethylhexyl)-phthalate in babies. Bio Neonate. 2000;78:269.

33. Bloom BT, Nelson RA, Dirksen HC. A new technique: umbilical arterial catheter placement. J Perinatol. 1986;6:174.

34. Squire SJ, Hornung TL, Kirchhoff KT. Comparing two methods of umbilical artery catheter placement. Am J Perinatol. 1990;7:8.

35. Stewart DL, Wilkerson S, Fortunate SJ. New technique for stabilizing umbilical artery catheters in very low birth weight infants. J Perinatol. 1989;9:458.

36. Clark JM, Jung AL. Umbilical artery catheterization by a cutdown procedure. Pediatrics. 1977;59:1036.37. Sherman NJ. Umbilical artery cutdown. J Pediatr Surg. 1977;12:723.

38. Hashimoto T, Togari H, Yura J. Umbilical artery cutdown: an improved procedure for reinsertion. Br J Surg. 1985;72:194.

39. Waffarn F, Devaskar UP, Hodgman JE. Vesico-umbilical fistula: a complication of umbilical artery cutdown. J Pediatr Surg. 1980;15:211.

40. Hepworth RC, Milstein JM. The transected urachus: an unusual cause of neonatal ascites. Pediatrics. 1984;73:397.

41. Dmochowski RR, Crandell SS, Corriere JN Jr. Bladder injury and uroascites from umbilical artery catheterization. Pediatrics. 1986;77:421.

42. Diamond DA, Ford C. Neonatal bladder rupture: a complication of umbilical artery catheterization. J Urol. 1989; 142:1543.

43. Kaufman JM, Sharada P, Austin TL, Macpherson RI. Neonatal bladder injury occurring after umbilical artery catheterization by cutdown. JAMA. 1983;250:2968.

44. Fakhraee SH, Krauss DR. Rupture of urinary bladder as a complication of umbilical artery cutdown. Clin Pediatr. 1979; 18:283.

45. Nagarajan VP. Neonatal bladder injury after umbilical artery catheterization by cutdown. JAMA. 1984;252:765.

46. Lehmiller DJ, Kanto WP Jr. Relationships of mesenteric thromboembolism, oral feeding and necrotizing enterocolitis. J Pediatr. 1978;92:96.

47. Davies MW, Mehr S, Morley CJ. The effect of draw-up volume on the accuracy of electrolyte measurements from neonatal arterial lines. J Pediatr Child Health. 2000;36:122.

48. Schulz G, Keller E, Haensse D, et al. Slow blood sampling from an umbilical artery catheter prevents a decrease in cerebral oxygenation in the preterm newborn. Pediatrics. 2003;111:e73.

49. Hermansen MC, Hermansen MG. Intravascular catheter complications in the neonatal intensive care unit. Clin Perinatol. 2005;32:141.

50. Miller D, Kirkpatrick BV, Kodroff M, et al. Pelvic exsanguination following umbilical artery catheterization in neonates. J Pediatr Surg. 1979;14:264.

51. Rao H, Elhassani S. Primum non nocere: iatrogenic complications of procedures performed on newborn. Part I: intravascular procedures. Perinatol Neonatol. 1980;4:25.

52. Vidyasager D, Downes JJ, Boggs TR. Respiratory distress syndrome of newborn infants: technique of catheterization of umbilical arteries and clinical results of treatment of 124 patients. Clin Pediatr. 1970;9:332.

53. Westrom G. Umbilical artery catheterization in newborns. V: a clinical follow-up study. Acta Paediatr Scand. 1980;69:371.

54. Macdonald MG, Chou MM. Preventing complications from lines and tubes. Semin Perinatol. 1986;10:224.

55. Stringel G, Mercer S, Richler M, et al. Catheterization of the umbilical artery in neonates: surgical implications. Can J Surg. 1985;28L:143.

56. Schreiber MD, Perez CA, Kitterman JA. A double-catheter technique for caudally misdirected umbilical arterial catheters. J Pediatr. 1984;104:768.

57. Chidi CC, King DR, Bates E. An ultrastructural study of intimal injury induced by an indwelling umbilical artery catheter. J Pediatr Surg. 1983;18:109.

58. Clark JM, Jung AL. Umbilical artery catheterization by a cut down procedure. Pediatrics (Neonatol Suppl). 1977;59:1036.

59. Lemons JA, Honeyfield PR. Umbilical artery catheterization. Perinat Care. 1978;2:17.

60. Hillman LS, Goodwin SL, Sherman WR. Identification of plasticizer in neonatal tissues after umbilical catheters and blood products. N Engl J Med. 1975;292:381.

61. James LS. Complications arising from catheterization of the umbilical vessels. In: 59th Ross Conference on Pediatric Research. Columbus, OH: Ross Laboratories; 1969:36.

62. Nagel JW, Sims SJ, Aplin CE II, et al. Refractory hypoglycemia associated with a malpositioned umbilical artery catheter. Pediatrics. 1979;64:315.

63. Urbach J, Kaplan M, Blondheim O, et al. Neonatal hypoglycemia related to umbilical artery catheter malposition. J Pediatr. 1985;106:825.

64. Carey BE, Zeilinger TC. Hypoglycemia due to high positioning of umbilical artery catheters. J Perinatol. 1989;9:407.

65. Van Leeuwen G, Patney M. Complications of umbilical artery catheterization: peritoneal perforation. Pediatrics. 1969;44:1028.

66. Malloy MH, Nichols MM. False abdominal aortic aneurysm: an unusual complication of umbilical arterial catheterization for exchange transfusion. J Pediatr. 1977;90:285.

67. Wynn ML, Rowen M, Rucker RW, et al. Pseudoaneurysm of the thoracic aorta: a late complication of umbilical artery catheterization. Ann Thorac Surg. 1982;34:186.

68. Wind ES, Wisoff G, Baron MG, et al. Mycotic aneurysm in infancy: a complication of umbilical artery catheterization. J Pediatr Surg. 1982;17:324.

69. Drucker DE, Greenfield LF, Salzberg AM. Aortic-iliac aneurysms following umbilical artery catheterization. J Pediatr Surg. 1986;21:725.

70. Lally KP, Sherman NJ. Iliac artery psuedoaneurysm following umbilical artery catheterization. Surgery. 1987; 101:636.

71. Kirpekar M, Augenstein H, Abiri M. Sequential development of multiple aortic aneurysms in a neonate post umbilical arterial catheter insertion. Pediatr Radiol. 1989;19:452.

72. Fok TF, Ha MH, Leung KW, et al. Sciatic nerve palsy complicating umbilical artery catheterization. Eur J Paediatr. 1986; 145:308.

73. Lynch MD. Sciatic palsy after umbilical artery catheterization. Brief report. J Bone Joint Surg (Br). 1988;70:151.

74. Himmel PD, Sumner DS, Mongkolsmai C, et al. Neonatal thrombosis associated with the umbilical arterial catheter: successful management by transaortic thrombectomy. J Vasc Surg. 1986;4:119.

75. Malin SW, Baumgart S, Rosenberg HK, et al. Nonsurgical management of obstructive aortic thrombosis complicated by renovascular hypertension in the neonate. J Pediatr. 1985; 106:630.

76. Seibert JJ, Northington FJ, Miers JF, et al. Aortic thrombosis after umbilical artery catheterization in neonates: prevalence of complications on long-term follow-up. AJR. 1991;156:567.

77. Martin JE, Moran JF, Cook LS, et al. Neonatal aortic thrombosis complicating umbilical artery catheterization: successful treatment with retroperitoneal aortic thrombectomy. Surgery. 1989;105:793.
78. Boo NY, Wong NC, Zulkifli SS, et al. Risk factors associated with umbilical vascular catheter-associated thrombosis in newborn infants. J Pediatr Child Health. 1999;35:460.

79. Cochrane WD, Davis HT, Smith CA. Advantages and complications of umbilical artery catheterization in the newborn. Pediatrics. 1968;42:769.

80. Egan EA, Eitzman DV. Umbilical vessel catheterization. Am J Dis Child. 1971;121:213.

81. De Sanctis N, Cardillo G, Nunziata Raga A. Gluteoperineal gangrene and sciatic nerve palsy after umbilical vessel injection. Clin Orthop. 1995;316:180.

82. Wigger HJ, Bransilver BR, Blanc WA. Thromboses due to catheterization in infants and children. J Pediatr. 1970;76:1.

83. Hermansen MC, Hermansen MG. Intravascular catheter complications in the neonatal intensive care unit. Clin Perinatol. 2005;32:141.

84. Cole ARF, Rolgin SH. A technique for rapid catheterization of the umbilical artery. Anesthesiology. 1980;53:254.

85. Dorand RD, Cook LN, Andrew BF. Umbilical vessel catheterization. The low incidence of complications in a series of 200 newborn infants. Clin Pediatr. 1977;16:569.

86. Gupta JM, Roberton NRC, Wigglesworth JS. Umbilical artery catheterization in the newborn. Arch Dis Child. 1968; 43:382.

87. Kitterman JA, Phibbs RH, Tooley WH. Catheterization of umbilical vessels in newborn infants. Pediatr Clin North Am. 1970;17:895.

88. Lividatis A, Wallgren G, Faxelius G. Necrotizing enterocolitis after catheterization of the umbilical vessels. Acta Paediatr Scand. 1970;63:277.

89. Bauer SB, Feldman SM, Gellis SS, et al. Neonatal hypertension: a complication of umbilical artery catheterization. N Engl J Med. 1975;293:1032.

90. Ford KT, Teplick SK, Clark RE. Renal artery embolism causing neonatal hypertension: a complication of umbilical artery catheterization. Radiology. 1974;113:169.

91. Plumer LB, Kaplan GW, Mendoza SA. Hypertension in infants: a complication of umbilical artery catheterization. J Pediatr. 1976;89:802.

92. Stevens PS, Mandell J. Urologic complications of neonatal umbilical artery catheterization. J Urol. 1978;120:605.

93. Butt WW, Gow R, Whyte H, et al. Complications resulting from use of arterial catheters: retrograde flow and rapid elevation in blood pressure. Pediatrics. 1985;76:250.

94. Brooks WG, Weibling RE. Emergency department presentation of severe hypertension secondary to complications of umbilical artery catheterization. Pediatr Emerg Care. 1987;3:104.

95. Krueger TC, Neblett WW, O'Neill JA, et al. Management of aortic thrombosis secondary to umbilical artery catheters in neonates. J Pediatr Surg. 1985;20:328.

96. Aziz EM, Robertson AF. Paraplegia: a complication of umbilical artery catheterization. J Pediatr. 1973;82:1050.

97. Lackey DA, Taber P. An unusual complication of umbilical artery catheterization. Pediatrics. 1972;49:281.

98. Haldman S, Fawter GW, Ashwal S, et al. Acute flaccid neonatal paraplegia: a case report. Neurology. 1983;33:93.

99. Brown MS, Phibbs RH. Spinal cord injury in newborns from use of umbilical artery. J Perinatol. 1988;8:105.

100. Henry CG, Gutierrez F, Joseph I, et al. Aortic thrombosis presenting as congestive heart failure: an umbilical artery catheter complication. J Pediatr. 1981;98:820.

101. Krishnamoorthy KS, Fernandez J, Todres ID, et al. Paraplegia associated with umbilical artery catheterization in the newborn. Pediatrics. 1976;58:443.

102. Martin GJ, Ireland WR. Letter to the editor: broken umbilical catheters. Am J Dis Child. 1977;131:1405.

103. Wagner CW, Vinocur CD, Weintraub WH. Retrieval of an umbilical artery catheter: a potential for misadventure. South Med J. 1987;80:1434.

104. Gaylord MS, Pittman PA, Bartness J, et al. Release of benzalkonium chloride from a heparin-bonded umbilical catheter with resultant factitious hypernatremia and hyperkalemia. Pediatrics. 1991;87:631.

105. Cochrane WD. Umbilical artery catheterization. In: Iatrogenic Problems in Neonatal Intensive Care. Report of the 69th Ross Conference of Pediatric Research. Columbus, OH: Ross Laboratories; 1976:28. 106. Hilliard J, Schreiner RL, Priest J. Hemoperitoneum associated with exchange transfusion through an umbilical arterial catheter. Am J Dis Child. 1979;133:216.

107. Sasidhanan P. Umbilical arterial rupture: a major complication of catheterization. Indiana Med. 1985;78:34.

108. Moncino MD, Kurtzberg J. Accidental heparinization in the newborn: a case report and brief review of the literature. J Perinatol. 1990;10:399.

109. Knudsen FV, Petersen S. Neonatal septic osteoarthritis due to umbilical artery catheterization. Acta Paediatr Scand. 1977; 66:225.

110. Krauss AN, Albert RF, Kannan MM. Contamination of umbilical catheters in the newborn infant. J Pediatr. 1970;77:965.

111. Larroche JCL, Bennoun M, Korn G. Umbilical catheterization: its complications (anatomical study). Symposium on Artificial Ventilation. Biol Neonate. 1970;16:101.

112. Lim MO, Gresham EL, Franken EA. Osteomyelitis as a complication of umbilical artery catheterization. Am J Dis Child. 1977;131:142.

113. Powers WF, Tooley WH. Letter to the editor: contamination of umbilical vessel catheters. Encouraging information. Pediatrics. 1968;49:470.

114. White AA II, Creline ES, McIntosh S. Septic arthritis of the hip joint secondary to umbilical artery catheterization associated with transient femoral and sciatic neuropathy. Clin Orthop. 1974;100:190.

115. Ruderman JW, Morgan MA, Klein AH. Quantitative blood cultures in the diagnosis of sepsis in infants with umbilical and Broviac catheters. J Pediatr. 1988;112:798.

116. Pourcyrous M, Korones SB, Bada HS, et al. Indwelling umbilical arterial catheter: a preferred sampling site for blood culture. Pediatrics. 1988;81:821.

117. Landers S, Moise AA, Fraley JK, et al. Factors associated with umbilical catheter related sepsis in neonates. Am J Dis Child. 1991;145:675.

118. Narendran V, Gupta G, Todd DA, et al. Bacterial colonization of indwelling vascular catheters in newborn infants. J Paediatr Child Health. 1996;32:391.

119. Castelman B, Scully RE, McNeely BV. Case records of the Massachusetts General Hospital. N Engl J Med. 1973; 189:1027.

120. Book LS, Herbst JJ. Intraarterial infusions and intestinal necrosis in the rabbit: potential hazards of umbilical artery injections of ampicillin, glucose, and sodium bicarbonate. Pediatrics. 1980;65:114.

121. Simpson JS. Misdiagnosis complicating umbilical vessel catheterization. Clin Pediatr. 1977;16:569.

122. Biagtan J, Rosenfeld W, Salazard D, et al. Herniation of the appendix through the umbilical ring following umbilical artery catheterization. J Pediatr Surg. 1980;15:672.

123. Bavikatte K, Hillard J, Schreiner RL, et al. Systemic vascular cotton fiber emboli in the neonate. J Pediatr. 1979;95:61.

124. Abramowsky CR, Chrenka B, Fanaroff A. Wharton jelly embolism: an unusual complication of umbilical catheterization. J Pediatr. 1980;96:739.

125. Diamond DA, Ford C. Neonatal bladder rupture: complication of umbilical artery catheteriztion. J Urol. 1989;142:1543.

126. Mata JA, Livne PM, Gibbons MD. Urinary ascites: complication of umbilical artery catheterization. Urology. 1987;30:375.

127. Dmochowski RR, Crandell SS, Lowiere JN. Bladder injury and uroascites from umbilical artery catheterization. Pediatrics. 1986;77:421.

128. McGravey VJ, Dabiri C, Bean MS. An unusual twist to umbilical artery catheterization. Clin Pediatr. 1983;22:587.

129. Rodriquez M, Sosenko I. Catheter-induced aortic thrombosis masquerading as coarctation of the aorta. Clin Pediatr (Phila). 1989;28:581.

130. Adelman RD, Morrell RE. Coarctation of the abdominal aorta and renal artery stenosis related to an umbilical artery catheter placement in a neonate. Pediatrics. 2000;106:E36.

131. Crie JS, Hajar R, Folger G. Umbilical catheter masquerading at echocardiography as a left atrial mass. Clin Cardiol. 1989; 12:728.

132. Sakurai M, Donnelly LF, Klosterman LA, et al. Congenital diaphragmatic hernia in neonates: variations in umbilical catheter and enteric tube position. Radiology. 2000;216:112.

29-Umbilical Vein Catheterization

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A. Indications

- Primary
 - \circ $\;$ Emergency vascular access for fluid and medication infusion and for blood drawing
 - Central venous pressure monitoring (if catheter across ductus venosus)
 - Exchange transfusion
- Secondary
 - Long-term central venous access in low-birthweight infants. If the line is to be used long-term, particularly if parenteral nutrition is to be infused by this route, the same aseptic techniques must be used to prevent line-related sepsis as are used for any central venous line. See Chapter 31.
 - Diagnosis of total anomalous pulmonary venous drainage below the diaphragm (1)

B. Contraindications

- Omphalitis
- Omphalocele
- Necrotizing enterocolitis
- Peritonitis

C. Equipment

- Catheter same as for umbilical artery catheterization, except:
 - 5-French (Fr) catheter for infants weighing <3.5 kg
 - 8-Fr catheter for infants weighing >3.5 kg
 - Catheter used for exchange transfusion (removed after procedure) should have side holes. This reduces risk of sucking thin wall of inferior vena cava against catheter tip, with possible vascular perforation (2).
- Other equipment as for umbilical artery catheter, but omit 2% lidocaine (see Chapter 28, C)

D. Precautions

- Keep catheter tip away from origin of hepatic vessels, portal vein, and foramen ovale. Catheter tip should lie in ductus venosus or inferior vena cava (3). Sometimes it will not be possible to advance the catheter through the ductus venosus. Vigorous attempts to advance are to be avoided. In an emergency, vital infusions (avoid very hypertonic solutions) may be given slowly after pulling catheter back into umbilical vein (approximately 2 cm) and checking blood return.
- Check catheter position prior to exchange transfusion. Avoid performing exchange transfusion with catheter tip in portal system or intrahepatic venous branch (see F.4).
- Once secured, do not advance catheter into vein.
- Avoid infusion of hypertonic solutions when catheter tip is not in inferior vena cava.
- Do not leave catheter open to atmosphere (danger of air embolus).
- Avoid use of central venous pressure monitoring catheter for concomitant infusion of parenteral nutrition (risk of sepsis).
- Be aware of potential inaccuracies of venous pressure measurements in inferior vena cava (see Chapter 31).

E. Technique (See Procedures DVD for Video)

Anatomic note: In the full-term infant, the umbilical vein is 2 to 3 cm in length and 4 to 5 mm in diameter. From the umbilicus, it passes cephalad and a little to the right, where it joins the left branch of the portal vein after giving off several large intrahepatic branches that are distributed directly to the liver tissue. The ductus venosus becomes a continuation of the umbilical vein by arising from the left branch of the portal vein, directly opposite where the umbilical vein joins it. At birth, it is 2 to 3 cm long and 4 to 5 mm in diameter, and it is located in a groove between the right and left lobes of the liver in the median sagittal plane of the body, at a level between the ninth and tenth thoracic vertebrae. It terminates in the inferior vena cava along with hepatic veins, as shown in Fig. 29.1.

• Make necessary measurements to determine length of catheter to be inserted, adding length of umbilical stump (Figs. 28.1 and 28.2) (4).



FIG. 29.1. Anatomy of the umbilical and associated veins, with reference to external landmarks.

- Prepare for procedure as for umbilical artery catheter (see Chapter 28, E).
- Identify thin-walled vein, close to periphery of umbilical stump (Fig. 29.2).
- Grasp cord stump with toothed forceps.
- Gently insert tips of iris forceps into lumen of vein and remove any clots.
- Introduce fluid-filled catheter, attached to the stopcock and syringe, 2 to 3 cm into vein (measuring from anterior abdominal wall).
- Apply gentle suction to syringe.
 - If there is not easy blood return, catheter may have a clot in tip. Withdraw catheter while maintaining gentle suction. Remove clot and reinsert catheter.
 - If there is smooth blood flow, continue to insert catheter for full estimated distance.



FIG. 29.2. The umbilical stump. Vein is indicated with an arrow.

- If catheter meets any obstruction prior to measured distance,
 - It has most commonly
 - Entered portal system, or
 - Wedged in an intrahepatic branch of umbilical vein
 - Withdraw catheter 2 to 3 cm, gently rotate, and reinsert in an attempt to get tip through ductus venosus.
- If the catheter is in the portal circulation, leave the misdirected catheter in its place. Pass a new 5-Fr catheter into the same vessel. Once the catheter is in good position, remove the misdirected catheter. This procedure has a success rate of 50% (5).
- Obtain radiographic verification of catheter position. A lateral radiograph is often necessary for exact localization (Fig. 29.3) (6, 7). Desired location is T9 to T10, just above the right diaphragm. Position of catheter tip may be estimated clinically by measurement of venous pressure (1) and observation of waveform (Figs. 29.4 and 29.5). The catheter has crossed the foramen ovale if the blood obtained is bright red (arterial in appearance). In this case, pull catheter back.
 - As soon as catheter has been advanced 2 to 3 cm into the vein, have an assistant connect to pressure-monitoring system (see Chapter 8).
 - While continuing to advance the catheter, measure venous pressure and note pressure changes with respiration (Fig. 29.4). Ideal position is with catheter tip in inferior vena cava, near right atrium, although placement in ductus venosus is acceptable for purposes other than measurement of central venous pressure.
- Secure catheter as for umbilical artery catheter (see Chapter 28, E).

There may be more bleeding from the umbilical vein than from the umbilical artery because the vein is not a contractile vessel. Local pressure is usually sufficient to stop oozing. For care of indwelling catheter, sampling technique, and removal of catheter, see Chapter 28.



FIG. 29.3. Anteroposterior (A) and lateral (B) radiographs demonstrating the normal course of an umbilical venous catheter, with an umbilical artery catheter (arrows) in position for comparison. Note how the venous catheter swings immediately superior from the umbilicus, slightly to the right as it traverses the ductus venosus into the inferior vena cava (IVC). The distal tip of this line is just superior to the right atrial–IVC junction, and it might optimally be pulled back slightly into the IVC. Note how the thinner umbilical artery catheter (arrows) heads inferiorly as it proceeds to the iliac artery and then ascends posteriorly and to the left until it reaches the level of D7.

B

F. Complications

- Infections (6, 8, 9, 10, 11, 12, 13, 14, 15, 16 and 17)
- Thromboembolic (8, 13, 18, 19)

Emboli from a venous catheter may be widely distributed. If catheter tip lies in the portal system and ductus venosus has closed, emboli will lodge in liver. If catheter has passed through ductus venosus, emboli will go to lungs; or because of right-to-left shunting of blood through foramen ovale or ductus arteriosus in sick newborn infants, emboli may be distributed throughout entire systemic circulation. These emboli may be infected and therefore may cause widespread abscesses.



FIG. 29.4. Venous and arterial pressure tracings may be used to facilitate placement and detect misplacement. A: The catheter has been pulled back through the ductus venosus, and the tip lies in the portal system. The portal venous pressure is higher than the central venous pressure, there are no venous pressure waves, and there is a small positive deflection during inspiration. B: Tip of catheter in the superior vena cava near the right atrium shows a deflection of more than 4 mm Hg during spontaneous inspiration (I) and a large negative deflection of more than 15 mm Hg during a sigh (S). Atrial tracing shows an AC and a V wave. AC wave occurs with atrial contraction and closure of atrioventricular valve after P wave of electrocardiogram. V wave occurs with ventricular contraction near T wave of electrocardiogram. (Based on data from Kitterman JA, Phibbs RH, Tooley WH. Catheterization of umbilical vessels in newborn infants. Pediatr Clin North Am. 1970;17:895, with permission.) C: Pressure tracing from right ventricle and pulmonary artery. Right ventricular pressure tracing shows a single large rise and fall, beginning just after onset of QRS complex. Pulmonary artery tracing usually shows a dicrotic notch at end of T wave. Diastolic pressure is higher than that in right ventricle. Pulmonary capillary wedge tracing should resemble atrial tracing, inasmuch as it reflects left atrial pressure transmitted to the catheter tip when anterograde pulmonary

arterial flow is occluded. Note: The marked negative deflection in the right atrial tracing would be more typically seen in infants who are receiving mechanical ventilation and thus have a positive airway pressure that exceeds ventricular filling pressures during each inspiration. In a spontaneously breathing neonate, positive airway pressure occurs only during expiration and never exceeds ventricular filling pressures. There are extremely small changes in cardiac pressures (i.e., on inspiration: right atrial [RA] mean pressure ↑, 1 mm Hg; left atrial [LA] mean pressure ↓ 1 mm Hg; on expiration: RA pressure ↓ 1 mm Hg; LA pressure ↑ 1 mm Hg) during the respiratory cycle as a result of changes in venous filling or preload. Right and left atrial pressures remain approximately equal in both inspiration and expiration (39).

- Catheter malpositioned in heart and great vessels (Figs. 29.5 and 29.6)
 - Pericardial effusion/cardiac tamponade (cardiac perforation) (3, 20, 21 and 22)
 - Cardiac arrhythmias (23)
 - Thrombotic endocarditis (24)
 - Hemorrhagic infarction of lung (7)
 - Hydrothorax (catheter lodged in or perforated pulmonary vein) (25)
- Catheter malpositioned in portal system
 - Necrotizing enterocolitis (26, 27)
 - Perforation of colon (28, 29)
 - Hepatic necrosis (thrombosis of hepatic veins or infusion of hypertonic or vasospastic solutions into liver tissues) (Fig. 29.7) (12, 13, 19, 30, 31)
 - Hepatic cyst (32)





descending aorta





FIG. 29.5. A: Radiograph showing venous catheter that has crossed the ductus arteriosus into the thoracic aorta. B: In this situation, the arterial pressure markings were not helpful because the presence of pulmonary hypertension in the patient rendered the tracings from the pulmonary artery and descending aorta virtually identical.

- Other
 - Perforation of peritoneum (33)
 - Obstruction of pulmonary venous return (in infant with anomalous pulmonary venous drainage) (1)
 - Plasticizer in tissues (34)
 - Portal hypertension (18, 24, 35, 36)
 - Electrical hazard (see Chapter 28, I.C.3) (2)
 - Fungal mass in right atrium (37)
 - Pseudomass in left atrium (38)
 - Digital ischemia (39)
 - Pneumopericardium (40)



FIG. 29.6. Spectrum of malpositions of umbilical venous catheters (UVCs) ($A\hat{a}\in C$). A: UVC in right portal vein with secondary air embolization into portal venous system. B: UVC in splenic vein. UAC catheter in good position with its tip at D7. C: UVC extending through heart into the superior vena cava. D, E: Spectrum of malpositions of UVCs. The anteroposterior film (D) shows an indeterminate position of the UVC. The right atrium, the right ventricle, and the left atrium are all possibilities.

The lateral film (E) shows its posterior position, confirming its presence in the left atrium. The lateral film is particularly important in making this distinction. Measurement of the PO₂ in blood from the catheter will be diagnostic of misplacement, unless the infant has severe persistent pulmonary hypertension or other cause of severe intracardiac shunting. Fâ \in I: Spectrum of malpositions of UVCs. Series of radiographs demonstrating various malpositions of a venous catheter: right pulmonary artery (F), left main pulmonary artery (G), main pulmonary artery (H), and right ventricle (I).



FIG. 29.7. A: Hepatic infarction (darkened areas on anterior aspect of liver) related to umbilical vein catheter. B: Section through inferior aspect of liver to show internal appearance of infarcted areas (arrow).

References

1. Nickerson BG, Sahn DJ, Goldberg SJ, et al. Hazards of inadvertent venous catheterization in a patient with anomalous pulmonary venous drainage: a case report. Pediatrics. 1979;63:929.

2. Kitterman JA, Phibbs RH, Tooley WH. Catheterization of umbilical vessels in newborn infants. Pediatr Clin North Am. 1970;17:895.

3. Johns AW, Kitchen WH, Leslie DW. Complications of umbilical vessel catheters. Med J Aust. 1972;2:810.

4. Nn P. Localization of the umbilical catheter by post-mortem measurement. Arch Dis Child. 1966;41:69.5. Mandel D, Mimouni FB, Littner Y, et al. Double catheter technique for misdirected umbilical vein catheter. J Peds. 2001;139:5.

6. Baker DH, Berdon WE, James LS. Proper localization of umbilical arterial and venous catheters by lateral roentgenograms. Pediatrics. 1969;43:34.

7. Weber AL, Deluce S, Shannan DL. Normal and abnormal position of the umbilical artery and venous catheter on the roentgenogram and review of complications. AJR. 1974;20:361.

8. Anagnostakis D, Kamba A, Petrochilou V, et al. Risk of infection associated with umbilical vein catheterization: a prospective study in 75 newborn infants. J Pediatr. 1975;86:759.

9. Balagtas RC, Bell CE, Edward LD, et al. Risk of local and systemic infections associated with umbilical vein catheterization: a prospective study in 86 newborn patients. Pediatrics. 1971;48:359.

10. Brans YW, Ceballos R, Cassady G. Umbilical catheters and hepatic abscesses. Pediatrics. 1974;53:264. 11. Centers for Disease Control. Guidelines for prevention of intravascular device-related infections, parts 1 and 2. Am J Infect Control. 1996;24:262.
12. Raad II, Luna M, Kaliel S-AM, et al. The relationship between the thrombotic and infectious complications of central venous catheters. JAMA. 1994;271:1014.

13. Raad II, Hohn DC, Gilbreath J, et al. Prevention of central venous catheter-related infections by using maximal sterile barrier precautions during insertion. Infect Control Hosp Epidemiol. 1994;15:231.

14. Tariq AA, Rudolph N, Levine EJ. Solitary hepatic abscess in a newborn infant: a sequel of umbilical vein catheterization and infusion of hypertonic glucose solutions. Clin Pediatr. 1977;16:577.

15. Williams JW, Rittenberry A, Dillard R, et al. Liver abscess in the newborn: complication of umbilical vein catheterization. Am J Dis Child. 1973;125:111.

16. Noel GJ, O'Loughlin JE, Edelson PJ. Neonatal staphylococcus epidermitis right sided endocarditis: description of five catheterized infants. Pediatrics. 1988;82:234.

17. Wilkins EG, Manning D, Roberts C, et al. Quantitative bacteriology of peripheral venous cannulae in neonates. J Hosp Infect. 1985;6:209.

18. Oski FA, Allen DM, Diamond LK. Portal hypertensionâ€"a complication of umbilical vein catheterization. Pediatrics. 1963;31:297.

19. Sarrut S, Alain J, Allison F. Early complications of umbilical vein perfusion in the premature infant. Arch Fr Pediatr. 1969;26:651.

20. Purohit DM, Levkoff AH. Pericardial effusion complicating umbilical venous catheterizations. Arch Dis Child. 1977;52:520.

21. Walker D, Pellet JR. Pericardial tamponade secondary to umbilical vein catheters. J Pediatr Surg. 1972;7:79.

22. Savani RC, Valentini RP, Mimouni F. Pericardial effusion as a complication of umbilical venous catheterization. J Perinatol. 1990;10:443.

23. Egan EA, Eitzman DV. Umbilical vessel catheterization. Am J Dis Child. 1971;121:213.

24. Symchych PS, Krauss AN, Winchester P. Endocarditis following intracardiac placement of umbilical venous catheters in neonates. J Pediatr. 1977;90:287.

25. Kulkarni PB, Dorand RD. Hydrothorax: a complication of intracardiac placement of umbilical venous catheters. J Pediatr. 1979;94:813.

26. Livaditis A, Wallgren G, Faxelius G. Necrotizing enterocolitis after catheterization of the umbilical vessels. Acta Paediatr Scand. 1974;63:277.

27. Shah KJ, Corkery JJ. Necrotizing enterocolitis following umbilical vein catheterization. Clin Radiol. 1978;29:295.

28. Friedman A, Abellera R, Lidsky I, et al. Perforation of the colon after exchange transfusion in the newborn. N Engl J Med. 1970;282:796.

29. Lucey JF. Colonic perforation after exchange transfusion. N Engl J Med. 1969;280:724.

30. Venkatavaman PS, Babcock DS, Tsang RC, et al. Hepatic injury: a possible complication of dopamine infusion through an inappropriately placed umbilical vein catheter. Am J Perinatol. 1984;1:351.

31. Richter E, Globl H, Hoethusen W, et al. Intrahepatic calcifications in infants following umbilical venous catheterization. Ann Radiol (Paris). 1984;27:117.

32. Levkoff AH, Macpherson RI. Intrahepatic encystment of umbilical vein catheter infusate. Pediatr Radiol. 1990;20:360.

33. Kanto WP, Parrish RA. Perforation of the peritoneum and intraabdominal hemorrhage. Am J Dis Child. 1977;131:1102.

34. Hillman LS, Goodwin SL, Sherwin WR. Identification and measurement of plasticizer in neonatal tissues after umbilical catheters and blood products. N Engl J Med. 1975;292:381.

35. Erkan V, Blankenship W, Stahlman MT. The complications of chronic umbilical vessel catheterization [abstr]. Pediatr Res. 1968;2:317.

36. Lauridsen UB, Enk B, Gammeltoft A. Oesophageal varices as a late complication of neonatal umbilical vein catheterization. Acta Paediatr Scand. 1978;67:633.

37. Johnson DE, Bass JL, Thomson TR, et al. Candida septicemia and right atrial mass secondary to umbilical vein catheterization. Am J Dis Child. 1981;135:275.

38. Crie JS, Hajar R, Folger G. Umbilical catheter masquerading at echocardiography as a left atrial mass. Clin Cardiol. 1989;12:728.

39. Welibae MA, Moore JH. Digital ischemia in the neonate following intravenous therapy. Pediatrics. 1985;76:99.40. Long WA. Pneumopericardium. In: Long WA, ed. Fetal and Neonatal Cardiology. Philadelphia: WB Saunders; 1990:382.

30-Peripheral Arterial Cannulation

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Arterial access is often needed in the care of the sick neonate. For various technical or clinical reasons, catheterization of the umbilical artery is not always possible. Therefore, peripheral arterial cannulation may be required. Although cannulation of the axillary (1,2) and temporal arteries (3,4) has been reported in neonates, these sites are not recommended because of the difficulty of access and potential adverse neurologic sequelae, respectively (5,6,7 and 8). Similarly, cannulation of the brachial artery should be avoided because of the poor collateral blood flow. As a general rule, the most peripheral available artery should be used, to reduce the potential sequelae from any associated vascular compromise or thromboembolic event. The artery chosen should be large enough to measure blood pressure without occlusion, have adequate collateral circulation, be at a site with low infection risk, and be in an area that can be easily monitored and cared for by nursing staff. Common sites for peripheral arterial cannulation include the radial, ulnar, dorsalis pedis, and posterior tibial arteries.

A. Indications

- Monitoring of arterial blood pressure
- Frequent monitoring of blood gases or laboratory tests (e.g., sick ventilated neonates or extremely low-birthweight premature infants)
- When preductal measurement is required (e.g., with persistent pulmonary hypertension) in the case of right-upper-extremity cannulation
- **B.** Contraindications
 - Bleeding disorder that cannot be corrected
 - Pre-existing evidence of circulatory insufficiency in limb being used for cannulation
 - Evidence of inadequate collateral flow (i.e., occlusion of the vessel to be catheterized may compromise perfusion of extremity)
 - Local skin infection
 - Malformation of the extremity being used for cannulation
 - Previous surgery in the area (especially cutdown)

C. Equipment Sterile

- Gloves
- Antiseptic solution (e.g., an iodophor)
- 4- x 4- in gauze squares
- 0.5 to 0.95 N saline with 1 to 2 U/mL heparin

Although hypernatremia has been reported in very small premature infants who received excess sodium in flush solution (9), in our experience 0.5 N saline has been used without complications at infusion rates of 0.5 to 1 mL/hr. Using heparinized saline has been shown to maintain line patency longer than hypotonic solutions such as heparinized 5% dextrose water (10) or unheparinized normal saline (11).

- 3- or 5-mL syringe
- 20-gauge venipuncture needle (if using larger-size 22-gauge cannula)
- Appropriate-size cannula: 22-gauge x 1- in (2.5-cm), 24-gauge x 0.75-in, or 24-gauge x 0.56-in tapered or nontapered cannula with stylet for larger to smaller neonates, respectively
- Antiseptic ointment (optional)
- Arterial pressure transducer and extension tubing (see Chapter 8)
- 5-0 nylon suture with curved needle (optional)
- Needle holder (optional)
- Suture scissors (optional)
- T connector primed with heparinized flush solution
- Transparent, semipermeable dressing

Nonsterile

• Equipment for transillumination (see Chapter 12) or Doppler ultrasound

Use of Doppler ultrasound for localization of the artery (12,13 and 14) and assessment of the adequacy of the palmar circulation has been described (15,16).

- ¹/₂-in, water-resistant adhesive tape
- Materials for forearm restraint (see Chapter 3) for radial or ulnar cannulation
- A constant-infusion pump capable of delivering flush solution at rate of 0.5 to 1 mL/hr against back pressure

Additional Equipment Required for Cutdown Procedure

- All equipment except mask must be sterile.
- Gown and mask
- 0.5% lidocaine hydrochloride in labeled 3-mL syringe
- 11 scalpel and holder
- Two curved mosquito hemostats
- Nerve hook
- 5-0 nylon suture

D. Precautions

- When performing radial artery cannulation, always check ulnar collateral circulation using the Allen test (see Chapter 14) (17) or Doppler ultrasound (15,16) prior to undertaking procedure.
- When performing dorsalis pedis or posterior tibial cannulation, a modified Allen test can be performed by raising the foot, occluding the dorsalis pedis and posterior tibial arteries, releasing pressure over one, and monitoring for tissue perfusion within 10 seconds, although this technique is less reliable than testing in the hand (18).
- When performing radial or ulnar cannulation, avoid excessive hyperextension of wrist, because this may result in occlusion of artery and a false-positive Allen test (19) and has been associated with median nerve conduction block (20).

- Leave all fingertips/toes exposed so that circulatory status may be monitored. Examine limb frequently for changes in perfusion.
- Never ligate artery.
- Take care not to introduce air bubbles into cannula while assembling infusion system or taking blood samples.
- Make sure that a continuous pressure waveform tracing is displayed on a monitor screen at all times.
- Be aware that the blood pressure measured in the lower extremity may be 5 to 20 mm Hg higher than in the upper extremity, and may be delayed by one tenth of a second (18).
- Do not administer a rapid bolus injection of fluid via line, because there is a danger of retrograde embolization of clot or air (21). Flush infusion after sampling should be:
 - Minimal volume (0.3 to 0.5 mL)
 - Injected slowly
- To reverse arteriospasm, see Chapter 33.
- Use cannula for sampling only; no fluids other than heparinized saline flush solution should be administered via cannula.
- Remove cannula at first indication of clot formation or circulatory compromise (e.g., dampening of waveform on monitor). Do not flush to remove clots.
- Inspect cannula insertion site at least daily.
 - If signs of cellulitis are present, remove the cannula and send the cannula tip for culture. Also, send a wound culture if there is inflammation at the cutdown site.
 - Obtain a blood culture from a peripheral site if signs of sepsis are present.
 - Inspect the area distal and proximal to the insertion site for blanching, redness, cyanosis, or changes in temperature or capillary refill time.
- Remove cannula as soon as indications no longer exist.

E. Technique

Standard technique for percutaneous arterial cannulation

- Choose a site for cannulation and secure the appropriate limb.
 - Radial artery: This is the most routine site for cannulation. The infant's forearm and hand can be transilluminated with the wrist in extension 45 to 60 degrees (Fig. 30.1), making sure that fingers are visible to monitor distal perfusion. The artery can be palpated proximally to the transverse crease on the palmar surface of the wrist, medial to the styloid process of the radius, and lateral to the flexor carpi radialis (Fig. 30.2).



FIG. 30.1. Transillumination of the radial artery.

- Ulnar artery: In a small number of infants, the ulnar artery may be more easily located than the radial artery (22). If an Allen test indicates that the collateral blood supply is adequate, the ulnar artery may be cannulated using the same method as for a radial artery. The ulnar artery runs along the palmar margin of the flexor carpi ulnaris, radial to the pisiform bone. Caution is necessary when cannulating the ulnar artery because it runs next to the ulnar nerve and is smaller in caliber than the radial artery (Fig. 30.2).
- Dorsalis pedis artery: The dorsalis pedis artery can be found in the dorsal midfoot between the first and second toes with the foot held in plantar flexion (Fig. 30.3). It should be noted that the vascular anatomy of the foot is variable and the dorsalis pedis artery may be absent in some patients (23), whereas it may provide the main blood supply to the toes in others (24).



FIG. 30.2. Anatomic relations of the major arteries of the wrist and hand.

- Posterior tibial artery: The posterior tibial artery runs posterior to the medial malleolus with the foot held in dorsiflexion (Fig. 30.4).
- Identify artery by
 - Palpation (see anatomic landmarks as described above or individual arterial sites)
 - Transillumination (Fig. 12.1 and Chapter 12) (25)
 - Doppler ultrasound (12,13 and 14)
- Scrub and put on gloves.
- Prepare skin over site with antiseptic (e.g., an iodophor).
- Make small skin puncture with venipuncture needle over site (to ease passage of cannula through skin and reduce chances of penetrating the posterior wall of the vessel, especially when using a larger-gauge cannula).
- Accomplish cannulation of artery (Fig. 30.5).
- Method A (preferred for small premature neonates) (Fig. 30.6)
 - Puncture artery directly at an angle of 10 to 15 degrees to the skin, with the needle bevel down.
 - Advance slowly. There will be arteriospasm when the vessel is touched, and blood return may be delayed.





FIG 30.3. A: Anatomic relations of dorsalis pedis artery. B: White arrow shows anatomic location of dorsalis pedis artery.

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FIG. 30.4. A: Anatomic relations of posterior tibial artery, showing site of incision for cutdown. B: Cannulation of posterior tibial artery; cannula is attached to a transducer for continuous blood pressure monitoring.



FIG. 30.5. A: Cannulation of artery using Method A (see text). B: Cannulation of artery using Method B (see text). (Redrawn from Filston HC, Johnson DG. Percutaneous venous cannulation in neonates and infants: a method for catheter insertion without "cutdown.†Pediatrics. 1971;48:896, with permission of American Academy of Pediatrics.)

• Withdraw needle stylet (blood should appear in the cannula) and advance cannula into artery as far as possible.

Method B (Fig. 30-5b)

- Pass needle stylet (with bevel up) and cannula through artery at 30- to 40-degree angle to skin.
- Remove stylet and withdraw cannula slowly until arterial flow is established.
- \circ Advance cannula into artery.

Inability to insert the cannula into the lumen usually indicates failure to puncture the artery centrally. This often results in laceration of the lateral wall of the artery with formation of a hematoma, which can be seen on transillumination.



FIG. 30.6. A: Puncture artery directly at angle of 10 to 15 degrees to skin, with needle bevel down. B: Advance slowly. C: Withdraw needle stylet, allow for blood return, and advance cannula into artery. D: Attach cannula firmly to T connector.

- Attach cannula firmly to T connector and gently flush with 0.5 mL of heparinized solution, observing for evidence of blanching or cyanosis.
- Apply iodophor ointment (optional) to puncture site.
- Suture cannula to skin with 5-0 nylon suture if desired.

This step may be omitted as long as cannula is securely taped (Fig. 26.4); use of sutures may produce a more unsightly scar.

- Secure cannula as done with peripheral intravenous line, shown in Fig. 26.4. Transparent semipermeable dressing may be used in place of tape to allow continuous visualization of skin entry site. Guarantee that all digits are visible for frequent inspection.
- Maintain patency by attaching T connector to extension tubing or arterial pressure line to run 0.5 to 1 mL/hr of heparinized flush solution by constant infusion pump.
- Change intravenous (IV) tubing and flushing solution every 24 hours.

Radial artery cutdown

- Cutdown technique is better for the very small neonate, because trauma to the artery causes vasospasm, which makes percutaneous cannulation of a small vessel very difficult.
- Technique I: Cutdown at wrist
- The artery is initially exposed by cutdown, and a catheter is inserted under direct vision.
 - Prepare as for percutaneous procedure (Standard Technique, Steps 1 to 3).
 - Scrub and prepare as for major procedure (see Chapter 4).

- Infiltrate site of incision (point of maximum pulsation just proximal to proximal wrist crease) with 0.5 to 1 mL of lidocaine.
- Wait 5 minutes for anesthesia.
- \circ Make a 0.5-cm transverse skin incision (Fig. 30.7A).
- Deepen incision into subcutaneous tissue by blunt longitudinal dissection with curved mosquito hemostat (Fig. 30.7B).
- Use curved mosquito hemostat to dissect artery free.

Be gentle, to avoid arteriospasm.

- Elevate artery with hemostat or nerve hook (Fig. 30.7C).
- Loop ligature (5-0 silk) around artery for traction purposes (Fig. 30.7D). Do not tie ligature.
- Advance cannula stylet into artery with bevel down, until cannula is clearly within vessel lumen (Fig. 30.7E).
- Remove stylet and advance cannula to hub (Fig. 30.7F).
- Remove ligature.
- See percutaneous method under E (Standard Technique, Steps 7 to 11) for fixation and care of cannula.

The incision can usually be kept small enough so that the hub of the cannula fills it and no closing suture is needed.

- Technique II: Cannulation at anatomic snuff box
 - Described by Amato et al. (26)
 - May be used in infants who have undergone previous arterial cutdown at wrist
 - Should not be a primary approach to radial artery (particularly if cannulation is achieved by cutdown)
 - Site is not easy to expose.
 - Scar tends to be more disfiguring than at wrist.
 - The radial artery passes dorsally at the wrist and traverses the anatomic snuff box, which is bounded medially by the extensor pollicus longus and extensor pollicus brevis muscles (Fig. 30.8A).
 - The artery becomes superficial immediately after passing the extensor pollicus longus and before passing beneath the first dorsal interosseous muscle.
 - The point for cannulation is located at the junction of a line drawn along the medial aspect of the extended thumb and another line drawn along the lateral aspect of the extended index finger (Fig. 30.8).

Posterior tibial artery cannulation by a cutdown procedure

- Prepare as for percutaneous method.
- Put on mask.
- Tape foot to footboard in equinovarus position (see Chapter 3).
- Scrub and prepare as for major procedure (see Chapter 4).
- Infiltrate incision site with 0.5 to 1 mL of 0.5% lidocaine (Fig. 30.4).
- Wait 5 minutes for anesthesia.
- Make transverse incision (0.5 cm) posteroinferior to medial malleolus (see Fig. 30.4).

A vertical, rather than a transverse, incision is optional. The former has the advantage that it offers the opportunity to extend the incision cephalad, should the posterior wall of the vein be perforated on the initial attempt at cannulation. However, it has the disadvantage that it may be made too far lateral or medial to the artery.

- Identify artery by longitudinal dissection with mosquito hemostat. The artery is usually found just anterior to the Achilles tendon and adjacent to the tibial nerve.
- Place mosquito hemostat behind artery, and loop 5-0 nylon suture loosely around it.

Be gentle, to avoid arteriospasm.

- Elevate artery in wound with suture. Do not ligate artery.
- While stabilizing artery with suture, insert needle and cannula, with bevel down.
- Withdraw stylet and advance cannula to hub.
- Remove nylon suture.
- Close wound with 5-0 nylon suture (usually requires only one suture).



FIG. 30.7. Radial artery cannulation by cutdown. A: Making transverse skin incision. B: Blunt dissection with mosquito hemostat. C: Elevating artery with artery hook. D: Looping ligature around artery. E: Introducing cannula into artery while gentle $\hat{a} \in \hat{c}$ back traction $\hat{a} \in \hat{c}$ is applied to suture. F: Cannula advanced to hub.



FIG. 30.8. A: Anatomic relations of the radial artery on the volar aspect of the wrist. B: Point for cannulation of the radial artery is indicated by the junction of the dotted lines. (Redrawn from Amato JJ, Solod E, Cleveland RJ. A second radial artery for monitoring the perioperative pediatric cardiac patient. J Pediatr Surg. 1977;12:715, with permission.)

• See percutaneous method under E (Standard Technique, Steps 7 to 11) for fixation and care of cannula.

F. Obtaining Arterial Samples Equipment

- Gloves
- Alcohol swabs
- Sterile 2- x 2-in gauze squares (for three-drop method)
- 25-gauge straight needle (for three-drop method)
- Appropriate-sized syringe for sample (heparinized if sample is not processed on site)
- Syringe with flush (for stopcock method)
- Ice if necessary for sample preservation
- Specimen labels and requisition slips

Technique I: Three-drop method

- Wash hands and put on gloves.
- Clean diaphragm of T connector with antiseptic solution and allow to dry.
- Clamp T-connector tubing close to hub.
- Place sterile gauze squares beneath hub.
- Introduce 25-gauge needle through diaphragm and allow 3 to 4 drops of fluid/blood to drip onto gauze.
- Attach syringe to needle and withdraw sample.
- Remove needle from diaphragm.
- Unclamp T connector and allow residual pump pressure to flush catheter (27,28).

Technique II: Stopcock method (a three-way stopcock needs to be interposed between the patient and the transducer)

- Wash hands and put on gloves.
- Clean hub of stopcock with antiseptic solution.
- Attach syringe to stopcock.
- Turn stopcock off toward infusion pump side.
- Aspirate waste (volume depends on length of tubing).
- Using second syringe, withdraw sample.
- Flush cannula slowly over 30 to 60 seconds with 0.5 mL of flush solution.
- Open stopcock to pump to allow for continued infusion of heparinized saline.

G. Removal of the Cannula

Indications

- Stabilization or resolution of the indications for cannulation of the artery
- Cannula-related infection
- Evidence of thrombosis or mechanical occlusion of the artery



FIG. 30.9. Complication of cannulation of the radial artery. Arrow indicates necrotic area on forearm.

Technique

- Remove tape/dressing and cut stitch (if present) securing cannula to skin.
- Remove cannula gently.
- Apply local pressure for 5 to 10 minutes.

H. Complications of Peripheral Arterial Cannulation

- Thromboembolism/vasospasm/thrombosis
 - Blanching of hand and partial loss of digits (29,30 and 31)
 - Gangrene of fingertips and hemiplegia (32)

- Necrosis of forearm and hand (Fig. 30.9) (33,34 and 35)
- Skin ulcers (34,36)
- Ischemia/necrosis of toes (Fig. 30.10)(22,37)
- Cerebral emboli (5,7,21,39)
- Reversible occlusion of artery (39,40)
- Infiltration of infusate (37)
- Infection (27,41,42,43,44,45 and 46)
- Hematoma (47,48)
- Damage to peripheral nerves, for example:
 - Median nerve above medial epicondyle of humerus (Fig. 30.11A) may affect the following:
 - Pronation of forearm
 - Abduction of wrist
 - Flexion of wrist and distal phalanges of middle and index fingers



FIG. 30.10. Complication of cannulation of dorsalis pedis artery. Healing areas of sloughed skin are seen at site of skin puncture on dorsum of foot and also on anterior aspect of lower leg. Tips of toes 1, 3, 4, and 5 are necrotic.



FIG. 30.11. A: Muscles supplied by the median nerve in the forearm. B: Muscles supplied by the median nerve in the hand. C: Muscles supplied by the ulnar nerve in the hand. D: Muscles supplied by the posterior tibial nerve in the ankle and foot.

- Opposition, abduction, and flexion of thumb (atrophy of thenar eminence)
- Sensationâ€"maximally over volar aspect index and middle fingers
- Vasomotor control in limb
- Median nerve at wrist (Fig. 30.11B) causes carpal tunnel syndrome (20,49).
- Ulnar nerve at wrist causes (Fig. 30.11C):

0

0

• Atrophy of small hand muscles

- Sensory loss over dorsal and palmar surfaces of ring and little fingers and ulnar portion of hand and wrist
- Peripheral portion of deep peroneal nerveâ€"anesthesia of the lateral aspect of the dorsum of the hand, which results in no significant disability
- Posterior tibial nerve at medial malleolus (Fig. 30.11D) may affect:
 - Flexor hallucis brevis muscle
 - Flexor of proximal phalanx of big toe
 - Muscles of foot that spread and close toes and flex proximal phalanx of toes
 - Sensation on plantar surface of foot

Lesions of posterior tibial nerve may be difficult to detect on examination but may lead to significant discomfort in later life owing to loss of plantar arches on weight bearing.

- False cortical thumbs (50)
- Burns from transilluminator (51,52)
- Hemorrhage (including accidental dislodgement of cannula) (37,39,53,54)
- Hypernatremia caused by heparinized saline infusion through cannula (9)
- Hypervolemia related to continuous flush device (55)
- Air embolism (38,52)
- Pseudoaneurysm (44)
- Acquired bone dysplasia (56)

References

1. Greenwald BM, Notterman DA, Debruin WJ, et al. Percutaneous axillary artery catheterization in critically ill infants and children. J Pediatr. 1990;117:442.

2. Lawless S, Orr R. Axillary monitoring of pediatric patients. Pediatrics. 1989;84:273.

3. Au-Yeung JB, Sugg VM, Kantor NM, et al. Percutaneous catheterization of scalp arteries in sick infants. J Pediatr. 1977;91:106.

4. Prian GW. Temporal artery catheterization for arterial access in the high risk newborn. Surgery. 1977;82:734.

5. Au-Yeung JB, Sugg VM, Kantor NM, et al. Letter to the editor: sequelae of temporal artery catheterization. J Pediatr. 1978;93:895.

6. Bull MJ, Schreiner RL, Bhuwan PG, et al. Neurologic complications following temporal artery catheterization. J Pediatr. 1980;96:1071.

7. Prian GW, Wright GB, Rumach CM, et al. Apparent cerebral embolization after temporal artery catheterization. J Pediatr. 1978;93:115.

8. Simmons MA, Levine RL, Lubchenco LO, et al. Warning: serious sequelae of temporal artery catheterization. J Pediatr. 1978;92:284.

9. Hayden WR. Hypernatremia due to heparinized saline infusion through a radial artery catheter in a very low birth weight infant. J Pediatr. 1978;92:1025.

10. Rais-Bahrami K, Karna P, Dolanski EA. Effect of fluids on life span of peripheral arterial lines. Am J Perinatol. 1990;7:122.

11. Clifton GD, Branson P, Kelly HJ, et al. Comparison of NS and heparin solutions for maintenance of arterial catheter patency. Heart Lung. 1991;20:316.

12. Buakham C, Kim JM. Cannulation of a nonpalpable artery with the aid of a Doppler monitor. Anesth Analg. 1977;56:125.

13. Nagabhushan S, Colella JJ, Wagner R. Use of Doppler ultrasound in performing percutaneous cannulation of the radial artery. Crit Care Med. 1976;4:327.

14. Maher JJ, Dougherty JM. Radial artery cannulation guided by Doppler ultrasound. Am J Emerg Med. 1989;7:260.

15. Mozersky DJ, Buckley CJ, Haywood CO Jr, et al. Ultrasound evaluation of the palmar circulation: a useful adjunct to radial artery cannulation. Am J Surg. 1973;126:810.

16. Morray JP, Brandford HG, Barnes LF, et al. Doppler-assisted radial artery cannulation in infants and children. Anesth Analg. 1984;63:346.

17. Allen EV. Thromboangiitis obliterans: methods of diagnosis of chronic occlusive arterial lesions distal to the wrist with illustrative cases. Am J Med Sci. 1929;178:237.

 Johnstone RE, Greenhow DE. Catheterization of the dorsalis pedis artery. Anesthesiology. 1973;39:654.
 Greenhow DE. Incorrect performance of Allen's test: ulnar artery flow erroneously presumed inadequate. Anesthesiology. 1972;37:356.

20. Chowet AL, Lopez JR, Brock-Utne JG, et al. Wrist hyperextension leads to median nerve conduction block: implications for intra-arterial catheter placement. Anesthesiology. 2004;100:287.

21. Loewenstein E, Little JW, Hing HC. Prevention of cerebral embolization from flushing radial artery cannulae. N Engl J Med. 1971;285:1414.

22. Barr PA, Summers J, Wirtshafter D, et al. Percutaneous peripheral arterial cannulation in the neonate. Pediatrics (Neonatol Suppl). 1977;59:1058.

23. Huber JF. The arterial network supplying the dorsum of the foot. Anat Rec. 1941;80:373.

24. Spoerel WE, Deimling P, Aitkin R. Direct arterial pressure monitoring from the dorsalis pedis artery. Can Anaesth Soc J. 1975;22:91.

25. Wall PM, Kuhns LR. Percutaneous arterial sampling using transillumination. Pediatrics. 1977;59:1032.
26. Amato JJ, Solod E, Cleveland RJ. A "second†radial artery for monitoring the perioperative pediatric cardiac patient. J Pediatr Surg. 1977;12:715.

27. Todres ID, Rogers MC, Shannon DE, et al. Percutaneous catheterization of the radial artery in the critically ill neonate. J Pediatr. 1975;87:273.

28. Stanford VF, Garcia-Prats JA, Adams JM. Radial artery catheters (RAC) in newborns: maintenance techniques and factors affecting duration [abstr]. Pediatr Res. 1982;16:310A.

29. Randel SN, Tsang BHL, Wung JT, et al. Experience with percutaneous indwelling percutaneous catheterization in neonates. Am J Dis Child. 1987;141:848.

30. Adams JM. Iatrogenic problems in neonatal intensive care: Report of the 69th Ross Conference on Pediatric Research. Columbus, OH: Ross Laboratories; 1976:32.

31. Cartwright GW, Schreiner RL. Major complications secondary to percutaneous radial artery catheterization in the neonate. Pediatrics. 1980;65:139.

32. Sammaan HA. The hazards of radial artery pressure monitoring. J Cardiovasc Surg (Torino). 1971;12:342.

33. Mayer T, Matlak ME, Thomson JA. Necrosis of the forearm following radial artery catheterization in a patient with Reye's syndrome. Pediatrics. 1980;65:141.

34. Wyatt R, Glaves I, Cooper DJ. Proximal skin necrosis after radial artery cannulation. Lancet. 1974;1:1135.

35. Hack WW, Vos A, Okken A. Incidence of forearm and hand ischemia related to radial artery cannulation in newborn infants. Intensive Care Med. 1990;16:50.

36. Ward RJ, Green HD. Arterial puncture as a safe diagnostic aid. Surgery. 1965;57:672.

37. Spahr RC, Macdonald HM, Holzman IR. Catheterization of the posterior tibial artery in the neonate. Am J Dis Child. 1979;133:945.

38. Chang C, Dughi J, Shitabata P, et al. Air embolism and the radial arterial line. Crit Care Med. 1988;16:141.

39. Miyasaka K, Edmonds JF, Conn AW. Complications of radial artery lines in the pediatric patient. Can Anaesth Soc J. 1976;23:9.

40. Hack WW, Vos A, Vanderlei J, et al. Incidence and duration of total occlusion of the radial artery in newborn infants after catheter removal. Eur J Pediatr. 1990;149:275.

41. Ducharme FM, Garthier M, Lacroix J, et al. Incidence of infection related to arterial catherterization in children: a prospective study. Crit Care Med. 1988;16:272.

42. Adams JM, Speer ME, Rudolph AJ. Bacterial colonization of radial artery catheters. Pediatrics. 1980;65:94.

43. Band JD, Maki DG. Infections caused by arterial catheters used for hemodynamic monitoring. Am J Med. 1979;67:735.

44. Cohen A, Reyes R, Kirk M, et al. Oster's nodes, pseudoaneurysm formation and sepsis complicating percutaneous radial artery cannulation. Crit Care Med. 1984;12:1078.

45. Ricard P, Martin R, Marcoux A. Protection of indwelling vascular catheters: incidence of bacterial contamination and catheter-related sepsis. Crit Care Med. 1985;13:541.

46. Galvis AG, Donahoo JS, White JJ. An improved technique for prolonged arterial catheterization in infants and children. Crit Care Med. 1976;4:166.

47. Adams JM, Rudolph AJ. The use of indwelling radial artery catheters in neonates. Pediatrics. 1975;55:261.

48. Brown AE, Sweeny DB, Tumley J. Percutaneous radial artery cannulation. Anesthesia. 1969;24:532.49. Koenigsberger MR, Moessinger AC. Iatrongenic carpal tunnel syndrome in the newborn infant. J Pediatr. 1977;91:443.

50. Skoglund RR, Giles EE. The false cortical thumb. Am J Dis Child. 1986;140:375.

51. Uy J, Kuhns LR, Wall PM, et al. Light filtration during transillumination. A method to reduce heat build-up in the skin. Pediatrics. 1977;60:308.

52. Stein RT, Kuhns LR. Letter to the editor. J Pediatr. 1978;93: 162.

53. Bedford RF, Wollman H. Complications of percutaneous radial artery cannulation: an objective perspective in man. Anesthesiology. 1973;38:228.

54. Downs JB, Rackstein AD, Klein EF, et al. Hazards of radial artery catheterization. Anesthesiology. 1973;38:283.

55. Morray J, Todd S. A hazard of continuous flush systems for vascular pressure monitoring in infants. Anesthesiology. 1983; 38:187.

56. Seibert JT, McCarthy RE, Alexander JE, et al. Acquired bone dysplasia secondary to catheter related complications in the neonate. Pediatr Radiol. 1986;16:43.

31-Central Venous Catheterization

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Central venous catheters provide stable intravenous (IV) access to sick or low-birthweight infants who need long-term IV nutrition or medications (1).

A percutaneous central venous catheter, also known as a peripherally inserted central catheter (PICC), is a soft, flexible catheter that is inserted into a peripheral vein and threaded into the central venous system. Central venous lines may be placed by surgical cutdown when percutaneous access is not possible. Totally implantable vascular access devices (ports) are rarely used in neonates and are thus not included in this chapter.

Regardless of the method employed to obtain secure and reliable venous access, the clinician should be familiar with the technique and anatomic considerations unique to the approach. Some form of analgesia and sedation is generally required, with general anesthesia being reserved for more complex access cases. The majority of venous access procedures in the critically ill neonate are performed at the bedside rather than in the operating room.

A. Common Indications

- Total parenteral nutrition
- Long-term IV medication administration

- Administration of hyperosmolar IV fluids or irritating medications that cannot be administered through peripheral IV cannulas.
- Fluid resuscitation
- Repetitive blood draws (usually a secondary indication; catheters are not usually inserted primarily for this indication in neonates; only larger-lumen catheters may be used for blood draws without risk of clotting).

B. Relative Contraindications

There are no absolute contraindications, as the clinical situation dictates the need for venous access.

- Skin infection at insertion site
- Uncorrected bleeding diathesis (not a contraindication for percutaneous catheters inserted in distal peripheral venous sites)
- Ongoing bacteremia or fungal infection (which may cause catheter colonization and infection)
- The patient can be treated adequately with peripheral IV access. Central venous catheters have significant risks of complications and must not be used when peripheral venous access is possible and adequate.

C. General Precautions

- Central venous catheterization must be performed by trained individuals.
- Obtain informed consent prior to performing the procedure.
- Plan ahead: Success with PICC placement is higher if the catheter is inserted electively before peripheral veins are "used up†by frequent cannulations.
- Infant should be on a cardiorespiratory monitor during the procedure.
- Follow the manufacturer's instructions for catheter use.
- Follow strict sterile technique, because sepsis is the most common complication.
- Never leave a catheter in a position where it does not easily and repeatedly withdraw blood during the insertion procedure, to ensure that the tip is not lodged against a blood vessel or cardiac wall (2).
- Always confirm the position of the catheter tip by radiography prior to using it. This is especially true for catheters placed at bedside.
- If the catheter is used for hyperalimentation:
 - If possible, avoid use for any other purpose (such as medication administration or transfusion), because catheter manipulation increases the risk of sepsis (3).
 - Avoid using a stopcock in the line (increased potential for infection).
 - If possible, the line should be cared for by specifically trained personnel. Central line teams have been shown to decrease the frequency of catheter-related infections (4).

TABLE 31.1 Vessels Amenable to Central Ve	enous Access
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Blood Vessel	Recommended Technique
Upper extremity: cephalic, basilic, median cubital, or	Percutaneous or surgical
axillary vein	May be done at bedside
Lower extremity: saphenous vein or femoral vein	Percutaneous or surgical
	May be done at bedside
Scalp vein	Percutaneous technique, amenable only to PICC
	lines
External jugular vein	Percutaneous or surgical
	May be done at bedside
Internal jugular vein	Surgical technique
Common facial vein	May be done at bedside
Subclavian vein	Percutaneous technique, may be done at bedside

D. Vessels Amenable to Central Venous Access

Table 31.1 lists the sites usually used for central venous catheterization in the newborn. When venous access is extremely difficult because of superior vena cava or inferior vena cava occlusion, alternative sites such as the azygos system or the hepatic vein may be used for prolonged access, but there is little information about these approaches in neonates (5, 6).

E. Position of Catheter Tip (Fig. 31.1)



FIG. 31.1. Chest radiograph with PICC tip in appropriate position, just above junction of superior vena cava and right atrium.

• The catheter should be placed in as large a vein as possible, ideally outside the heart, and parallel with the long axis of the vein such that the tip does not abut the vein or heart wall (7). The recommendations for appropriate position of a central venous catheter tip vary, but there is general agreement that the tip should not be within the right atrium (8, 9, 10, 11). However, one large retrospective audit of 2,186 catheters showed that catheters with their tips in the right atrium and not coiled were not associated with pericardial effusions (2).

Recommendations for appropriate position of the catheter tip are as follows.

- The catheter tip should be within the superior or inferior vena cava, outside the pericardial reflection, a distance estimated to be about 1 cm outside the cardiac silhouette in preterm neonates and 2 cm outside the cardiac silhouette in term neonates (8).
- When inserted from the upper extremity, the catheter tip should be in the superior vena cava, outside the cardiac reflection and above the T2 vertebra (12).
- When inserted from the lower extremity, the catheter tip should be above the L4/ L5 vertebrae or the iliac crest, but not in the heart (12).
- \circ The tip of the catheter should be at the junction of the vena cava and the right atrium (7, 13).
- Confirmation of catheter tip placement
 - The tip of the radio-opaque catheter is usually seen on a routine chest radiograph (Fig. 31.1), but there can be significant interobserver variability in assessing the position (14). Digital enhancement of the radiograph may improve visualization but does not prevent interobserver variability (15).

- Two radiographic views (anteroposterior and lateral) help to confirm that the catheter is in a central vein. This is particularly important for catheters placed in a lower extremity, where the catheter may inadvertently be in an ascending lumbar vein and may appear to be in good position on an anteroposterior view (16).
- The use of radio-opaque contrast improves localization of the catheter tip, particularly when the catheter is difficult to see on a standard radiograph (17, 18). A 0.5-mL aliquot of 0.9% saline is instilled into the catheter to check patency, followed by 0.5 mL of iohexol. The radiograph is taken, and the line is flushed again with 0.5 mL of 0.9% saline. With this technique, there is no need to inject the contrast material while the radiograph is being taken (17). Other authors recommend injecting a 2-mL bolus of nonionic water-soluble contrast medium into the catheter during radiographic exposure, but this technique results in a larger volume of iodine-containing contrast being injected (18).
- Ultrasonography may also be useful in localizing the catheter tip (19).
- Chest radiographs obtained for any reason should be scrutinized for appropriate catheter position. Routine weekly radiographs taken for this purpose do not appear to reduce the risk of complications (13).

F. Methods of Vascular Access

- Percutaneous technique
 - o Advantages
 - Simpler to perform and relatively rapid procedure
 - May not require sedation
 - Vessel is not ligated as in open cutdown methods
 - Decreased potential for wound infection/dehiscence complications
 - o Disadvantages
 - Beyond the initial insertion into the peripheral vein, further passage of the catheter into its final position is essentially a blind technique.
 - Smaller-caliber catheter may preclude use for blood transfusions
 - Potential for injury to adjacent anatomic structures (20)
- Cutdown or open surgical technique
 - Advantages
 - Allows for insertion of larger silicone catheter (3- or 4.2-French [Fr])
 - The catheters can be tunneled under the skin away from the venotomy site, so they can remain in place longer with a lower risk of infection.
 - \circ Disadvantages
 - Requires general anesthesia/intravenous sedation
 - Requires surgical incision
 - Vein is often ligated, so it cannot be reused in the future.
 - Potential for injury to adjacent anatomical structures
 - Increased potential for wound infection
 - An operating room is the ideal setting for the procedure, so risks of transport of critically ill neonates need to be taken into consideration.

G. Types of Central Venous Catheters

	I ADLE JI	.2 Catheter Materials
Type of Catheter	Advantages	Disadvantages
Silicone	Soft, pliable	May be more difficult to insert percutaneously
	Lower risk of vessel	Thrombosis reported
	perforation	Fragile material: less tolerance to pressure
	Reported to be	Poor tensile strength: can tear or rupture
	thromboresistant	May be less radio-opaque
Polyurethane	Easier to insert	Increased risk of vessel perforation during insertion
	percutaneously	Thrombosis reported
	Stiffer on insertion but	
	softens within body	
	Some catheters are more	
	radio-opaque	
	Tensile strength: more	
	tolerant to pressure	
	Reported to be	
	thromboresistant	
Polyethylene	Easier to insert	High degree of stiffness may increase vessel perforation
	Very high tensile strength	during insertion or throughout catheter dwell
Polyvinyl chloride	Easier to insert	May leach plasticizers into body
(PVC)	percutaneously	High incidence of thrombosis
	Stiff on insertion but softens	
	within body	

TADLE 21 2 Catheter Materials

- Catheter materials: See Table 31.2.
- Types of catheters
 - Percutaneous (PICC) catheters/introducers

PICC catheters and kits are available commercially. PICCs are generally made of silicone or polyurethane. Sizes include 1.2, 1.9, 2, and 3 Fr. Larger sizes are generally not used in the neonatal population. Most catheters are single-lumen. Recently, a 1.9-Fr double-lumen catheter has become available. Double-lumen catheters can decrease the need for maintaining concurrent IV access when more than one site is required. PICC introducers/needles are available in 19, 20, 22, and 24 gauge. Choice will depend on the size of the vein to be cannulated.

Surgically placed central venous catheters 0

Surgically placed central venous catheters for neonates are available in sizes 2.5, 2.7, 3, 4.2, and 5 Fr. Catheters are usually silicone or polyurethane, with tissue in-growth cuffs that adhere to the subcutaneous tract, anchoring the catheter. Recently, antimicrobial cuffs have become available. Most catheters are single-lumen, but a few manufacturers make doublelumen catheters in sizes 5 Fr and larger.

Percutaneous Central Venous Catheterization (See Procedure DVD for Video)

A. Insertion sites (Fig. 13.1)

- Antecubital veins: basilic and cephalic veins
- Scalp veins: temporal and posterior auricular veins

- Saphenous veins
- Axillary vein
- External jugular vein

Right-sided and basilic veins are preferred because of the shorter and more direct route to the central vein. The cephalic vein may be more difficult to thread to the central position because of narrowing of the vessel as it enters the deltopectoral groove and the acute angle at which it joins the subclavian vein. The axillary and external jugular veins are the last choices because they are close to arteries and nerves.

- **B.** Insertion Variations
 - Breakaway needle: Needle is inserted into the vein. Next, the catheter is advanced through the needle. The needle is then retracted, split, and removed (Fig. 31.2). Disadvantage: There is a potential for shearing or severing the catheter if it is retracted while the needle is in the vein.
 - Peel-away introducer: A needle introducer is used to place a small cannula or sheath into the vein. The needle is then removed and the catheter is threaded through the cannula. The introducer cannula or sheath is then retracted from the vein, split or "peeled†apart, and removed from the catheter (Fig. 31.3).
 - Intact cannula: This technique is now rarely used because most commercially available catheters have a hub and introducer needles. A regular IV cannula is used to obtain venous access. The needle is removed. The catheter is threaded through the cannula to its final position. The cannula is then retracted and slipped off the end of the "hubless†catheter. A blunt needle with hub is connected to the end of the catheter (Fig. 31.4). Disadvantage: The blunt needle attachment must be secured well, otherwise leakage can occur.
- C. Placement of PICC Using the Break-Away or Peel-Away Introducer Needle
 - Equipment

All equipment used, except the mask, head cover, and tape measure, must be sterile. Commercial kits contain many of the necessary items.

- Radio-opaque central venous catheter
- o Break-away or peel-away needle introducer
- Tourniquet (optional)
- o Drapes
- Forceps
- Gauze pads
- Skin prep: 10% povidone–iodine or 0.5% chlorhexidine solution (as per institutional policy)
- Transparent dressing
- Sterile tape strips
- Sterile heparinized saline solution (1 U of heparin/mL or per institutional policy)
- 5- to 10-mL syringe with needle
- Tape measure
- Sterile surgical gown, sterile gloves, mask, and head cover
- Preparation
 - Although anesthesia is not required, nonpharmacologic comfort measures, pain medication, or sedation should be provided as needed. A small dose of sedative or narcotic analgesic may be useful.
 - Gather supplies. Wash hands thoroughly.

- Identify appropriate vein for insertion (see D).
- Position infant to facilitate insertion (Table 31.3). Restrain infant; provide comfort measures.
- Measure approximate distance from the insertion site to the point where the catheter tip will be placed (Table 31.3).
- Don mask and cap, scrub hands, and don sterile surgical gown and gloves.
- Trimming of the catheter to appropriate size is based on unit policy and manufacturer recommendations. Catheter is fragile. Handle it with care: Do not clamp or suture, and do not stretch or apply tension to catheter.
- Utilizing sterile technique, flush catheter with heparinized saline solution, leaving syringe attached. A small-barreled syringe (such as a 1-mL syringe) may generate too much pressure, resulting in catheter rupture (21).
- Prepare sterile field: Place drape under extremity. Utilizing the prep solution, prepare the area at and around the insertion site, working outward in concentric circles. Allow the prep solution to dry. Repeat process with new gauze/prep solution. Drape prepared area, leaving insertion site exposed.



FIG. 31.2. Peripherally inserted central catheter using break-away needle technique. (From Gesco International, San Antonio, TX, USA, with permission.)

- Catheter insertion
 - Utilizing a break-away needle (Fig. 31.2)
 - Apply tourniquet (optional).
 - Providing slight skin traction, insert needle about 1 cm below the intended vein. Insert needle at a low angle (approximately 15 to 30 degrees).
 - When a flashback is obtained, advance the needle about 1/4 in at a lower angle to ensure that the whole bevel of the needle is within the vein.

• Using nontoothed iris forceps, gently grasp the catheter about 1 cm from its distal end and insert it into the introducer needle.



FIG. 31.3. Peripherally inserted central catheter using a peel-away cannula introducer. A: Perform venipuncture. When flashback of blood is noted, reduce angle and advance introducer sheath farther to ensure placement in the vein. B: Withdraw the introducer needle from the sheath. Note that the introducer sheath is supported to avoid displacement. C: Insert the catheter into the introducer sheath using fine nontoothed forceps. D: Withdraw the introducer sheath. Note that the catheter is stabilized by applying digital pressure to the vein distal to the introducer sheath. E: Remove the introducer sheath by splitting and peeling it away from the catheter. Complete catheter advancement to premeasured length. F: Aspirate catheter to check for blood return and flush with heparinized saline to ensure patency. (From Klein C. NeoPicc: The Neonatal and Pediatric Workshop Manual. San Antonio, TX: Klein Baker Medical, 1998, with permission.)

- Caution: Never advance the needle or retract the catheter after inserting it into the needle; the catheter may be severed by this action.
- If a tourniquet was applied, it should be loosened or removed prior to advancement.
- With small, gentle nudges, a few millimeters at a time, advance the catheter through the needle to a distance of about 5 or 6 cm into the vein.

• Once the catheter is successfully advanced to about 5 or 6 cm, withdraw the needle carefully.



FIG. 31.4. Use of a blunt scalp vein needle to form a hub for a silicone catheter. The plastic needle cover is used to stabilize the needle \hat{a} the function. A commercially available blunt needle adapter may be inserted and fixed in a similar manner.

- To withdraw the needle, stabilize the catheter by applying gentle pressure over the vein proximal to the needle. Slowly withdraw the introducer needle until it is completely away from the site.
- Break the introducer needle by splitting the wings, and then carefully peel it away from the catheter.
- Continue to advance the catheter into the vein to the premeasured length, by nudging it farther, a few millimeters at a time, using the fine forceps.
- Difficulties in advancing catheter: Gently massage the vein in the direction of blood flow, proximal to the insertion site, or gently flush the catheter intermittently with 0.5 to 1.0 mL of heparinized saline; reposition the arm or head.
- Aspirate to visualize blood return in the catheter, and flush with 0.5 to 1 mL of heparinized saline to clear the catheter.
- Verify length of catheter inserted and adjust as necessary.
- Apply gentle pressure on insertion site with gauze pad to stop any bleeding.
- Secure catheter at skin insertion site with sterile tape strips and cover with sterile gauze until radiographic confirmation of position.

Site of Insertion	Position of Baby	Measurement
Antecubital	Supine, abduct arm 90 degrees from trunk;	From planned insertion site,
veins	turn head toward insertion site to prevent	along venous pathway, to
	catheter from traveling cephalad through	suprasternal notch, to third
	ipsilateral jugular vein	RICS
Saphenous or	Supine for greater saphenous vein, prone for	From planned insertion site,
popliteal	small saphenous or popliteal; extend leg	along venous pathway, to
veins		xiphoid process
Scalp veins	Supine, turn head to side; may have to turn	Follow approximate venous
	head to midline during procedure to assit	pathway from planned insertion
	advancement of catheter	site near ear, to jugular vein,
		right SC joint, to third RICS
External	Supine, turn head to side; place roll under neckFrom planned insertion site, to	
jugular vein	to cause mild hyperextension	right SC joint, to third RICS
Axillary vein	Supine, externally rotate and abduct arm 120	From planned insertion site, to
	degrees, flex forearm and place baby's hand	right SC joint, to thirt RICS
	behind head; vein is found above artery	
	between medial side of humeral head and	
	small tuberosity of the humerus	1 66

TABLE 31.3 Patient Position and Measurement for PICC Insertion Position of Rahy Magguramont

PICC, peripherally inserted central catheter; RICS, right intercostal space; SC, sternoclavicular.

Utilizing a peel-away cannula/introducer sheath (Figs. 31.3 and 31.5) 0

The procedure is similar to using a break-away needle.

- Apply tourniquet (optional). •
- Providing slight skin traction, insert needle about 1 cm below the intended vein. Insert needle at a low angle (approximately 15 to 30 degrees).



FIG. 31.5. A: Venipuncture with peel-away cannula introducer. B: Withdraw the introducer needle from the sheath. Note that the introducer sheath is supported to avoid displacement. C: Insert the catheter into the introducer sheath using fine nontoothed forceps. D: Withdraw the introducer sheath. E: Remove the introducer sheath by splitting and peeling it away from the catheter. F: Transparent dressing on PICC catheter. Note that the excess catheter length has been coiled in place under the dressing.

- When flashback is obtained, advance the introducer sheath about 1/4 in at a lower angle to ensure that the tip is within the vein.
- Withdraw the introducer needle while leaving the introducer sheath in place. Successful venipuncture is indicated by the flow of blood from the end of the sheath. Apply gentle pressure over insertion site to minimize blood loss. If a tourniquet was applied, it may be loosened or removed at this stage.
- Caution: Do not reintroduce the needle into the introducer sheath if the venipuncture is unsuccessful. This could result in a sheared or severed sheath.
- Using nontoothed iris forceps, gently grasp the catheter about 1 cm from its distal end and insert it into the introducer sheath.
- With small, gentle strokes, a few millimeters at a time, advance the catheter to the premeasured length.
- Aspirate to visualize blood return, and flush with 0.5 to 1 mL of heparinized saline to clear the catheter.

- Stabilize the catheter by applying gentle pressure to the vein proximal to the insertion site, then withdraw the introducer cannula. Carefully split and peel away the introducer sheath from the catheter.
- Once the introducer sheath has been removed, verify catheter insertion length and adjust with forceps.
- With a gauze pad, apply gentle pressure to insertion site to stop any bleeding.
- Secure catheter at skin insertion site with sterile tape strips, and cover with sterile gauze until radiographic confirmation of position.

D. PICC Dressings (Figs. 31.5 and 31.6)

- Antimicrobial prep solutions should be removed from the skin with sterile water or saline and allowed to dry before placing dressing.
- To prevent migration of the catheter, secure it to the skin a few millimeters from the insertion site with a small piece of sterile tape.
- If the catheter has not been trimmed, loosely coil the excess length of catheter close to the insertion site and secure to the skin with more sterile tape. Ensure that there is no kinking or stretching of the catheter under the dressing.
- Some manufacturers caution against placing tape that contains a wire directly on the silicone catheter. If the catheter is trimmed to the appropriate length, it may not be necessary to place sterile tape on the catheter.
- Apply a semipermeable transparent dressing over the area surrounding the insertion site.
- Do not allow tapes or transparent dressing to extend around the extremity. The dressing will form a constricting tourniquet as the infant grows or if there is venous congestion.
- Place tape under the catheter hub and criss-cross it over the hub (chevron). Do not obscure visualization of the insertion site (Fig. 31.6).
- To prevent skin breakdown, a skin barrier can be placed under the hub (e.g., Duoderm [Convatec: Bristol-Myers Squibb, Princeton, NJ, USA] or soft gauze). Ensure that the hub is secured.



FIG. 31.6. Peripherally inserted central catheter dressing with trimmed catheter. No excess catheter is present externally. The silicone heart is anchored with a piece of tape, and a sterile transparent dressing is placed over the insertion site. With use of a "chevron†technique, another piece of tape is placed under the catheter extension, next to the silicone heart, and crossed over on top of the transparent dressing. (From Klein C. NeoPicc: The Neonatal and Pediatric Workshop Manual. San Antonio, TX: Klein Baker Medical, 1998, with permission.)

E. Dressing Changes

- Mild oozing of blood from the insertion site may occur for up to 24 hours. If oozing occurs, the initial dressing should be changed when it subsides. If oozing of blood is a problem, a small piece of thrombolytic foam can be applied over the insertion site and under the dressing for the first 24 hours after insertion.
- The catheter site dressing should be replaced when it becomes damp, soiled, or loose. Transparent dressings should be changed every 7 days except in those patients in whom the risk of dislodging the catheter outweighs the benefit of changing the dressing (22).
- Inspect catheter site carefully at each dressing change (Table 31.4).
- Catheter may be pulled back if it is too far in, prior to replacing dressing. Do not advance catheter, as the risk of contamination is high.
- Use sterile technique for dressing changes (mask, cap, and sterile gloves; sterile gown is optional).
- Prepare sterile field: Place drape under extremity. Utilizing the prep solution, prepare the skin at and around the insertion site, working outward in concentric circles. Allow the prep solution to dry. Repeat process with new gauze/prep solution. Drape prepared area, leaving insertion site exposed.
- Follow steps D1 through D8 to complete the PICC dressing change.

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TABLE 31.4 Examination of the Catheter Site		
Assessment	Comments	
Catheter:	Catheter length should be clearly documented. If external length has	
Note external catheter length	changed, get radiograph to assess where the catheter tip is located.	
	If the catheter is pulled out, cover site with occlusive dressing and measure	
	catheter length to assure that some of the catheter was not retained in the vessel.	
Assess for kinks, tension,	Kinks and tension can damage catheter. It is recommended that damaged	
damage	catheters be removed, but some manufacturers provide repair kits.	
Insertion site/surrounding skin:	Mild erythema and/or phlebitis is common after the catheter is inserted. If	
Erythema, drainage, bleeding,	condition is severe and/or is persistent, consider removing catheter.	
edema, phlebitis, skin	Mild oozing of blood should not persist longer than 24 h.	
breakdown	Edema may be due to venous stasis from lack of extremity movement,	
	constrictive dressings, thrombus, damage to internal structures, localized	
	infection, or infiltration of infusion into soft tissue.	
	Avoid skin breakdown by utilizing skin barriers underneath hub, removing	
	dressing adhesives with care, minimizing tape, and removing antiseptics	
	from skin before applying dressing.	
Drainage/leaking	Purulent drainage may be due an infectious process. Consider obtaining	
	blood cultures and/or removing the catheter.	
	Clear drainage may be indicative of infusion leakage. This may be due to	
	catheter occlusion, infiltration, or damage to catheter.	

F. PICC Care and Maintenance

- Evaluate appearance of the catheter and the tissue around the insertion site frequently.
- Change tubing according to unit policy. Utilize aseptic technique when changing tubing.
- To prevent contamination of the line, enter the PICC only when absolutely necessary. Maintain sterility at connection site when entering the line.
- Do not utilize the PICC for routine blood sampling.
- Prime volumes are usually less than 0.5 mL. Use a 5- to 10-mL syringe when needed to check catheter patency. Do not use force if resistance is encountered. A small-barreled syringe (such as a 1-mL syringe) may generate too much pressure, resulting in catheter rupture (21).
- Add 0.5 to 1 U of heparin/mL of intravenous fluids. The effectiveness of heparin in the prevention of PICC occlusion and catheter thrombosis is unclear at present (23).
- Administer a constant infusion of intravenous fluids at a rate of at least 1 mL/h. Follow the manufacturer's recommendations for maximum flow rates.
- Ensure that medications infused through the line are compatible with IV fluids or that the line is flushed, before and after medication administration.
- Packed red blood cell transfusions should be given through a PICC only in an emergency, as the small catheter size may cause occlusion or hemolysis when older blood is used (24). A peripheral intravenous cannula should be utilized for blood transfusions.
- Monitor quality indicators in order to identify and solve problems. Infection rates, catheter dwell times, patient outcomes, and rates of complications should be monitored (22).
- Remove catheter as soon as it is no longer medically necessary by slowly withdrawing it from insertion site. Clean insertion site with prep prior to withdrawing catheter. Hold pressure over site if bleeding is a problem. Remove prep from skin. Place a clean gauze dressing over site. Document length removed.

Placement of Central Venous Catheters by Surgical Cutdown

A. Types of Catheters

Silicone catheters are preferred because they are constructed of relatively inert materials, offer increased pliability, and are associated with lower rates of infection and thrombosis. These catheters are placed in a central vein, and the distal end is tunneled subcutaneously a short distance from the access site to an exit wound. The catheters usually have a single lumen with a Dacron cuff, which adheres to the subcutaneous tract, anchoring the catheter. Polyethylene catheters have a higher rate of infection and thrombolytic complications and are not recommended for long-term intravenous access.

B. Contraindications

In addition to the relative contraindications delineated earlier, the internal jugular vein should be avoided if the contralateral jugular vein has been catheterized previously, or if there is thrombosis of the jugular venous system on the opposite side.

C. Equipment Sterile

- Skin prep: Per institutional policy (e.g., 10% povidone–iodine, or 0.5% chlorhexidine solution)
- Gown and gloves
- Cup with antiseptic solution
- Sterile transparent aperture drape; four sterile towels to ensure a sterile operative field
- Four 4-4-in gauze squares
- Local anesthetic: 0.5% lidocaine HCl in labeled 3-mL syringe with 25-gauge venipuncture needle

Consider sedation and pain medication in addition to local anesthesia. Patients who are intubated may be given a sedative and muscle relaxant in addition to local anesthesia. When patients are taken to the operating room, general anesthesia is preferred.

- Catheter of choice
- Heparinized 0.25 N saline flush solution (1 U/mL) in 3-mL syringe
- 4-0 Polyglactin suture (Vicryl; Ethicon, Somerville, NJ, USA) and 5-0 nylon suture (black monofilament nylon) on cutting needles (see Appendix B)
- T connector connected with a sterile 3-mL syringe filled with heparinized saline
- No. 11 scalpel blade and holder
- Two small tissue retractors or self-retaining retractor
- Tissue forceps
- Fine vascular forceps
- Two small, curved mosquito hemostats
- Dissecting scissors
- 4-0 Vicryl suture on small, curved needle; 6-0 polypropylene on a tapered needle. This is used for a purse-string stitch as an alternative to ligation of the vessels.
- Needle holder
- Suture scissors
- Appropriate materials for occlusive dressing of choice

Nonsterile

- Cap and mask
- Roll of 4- \tilde{A} 4-in gauze
- Tape measure
- Adhesive tape

D. Techniques

In the neonate, the cervical veins are preferable to the lower-extremity veins. The cervical veins are easily accessible and are a proportionately larger size. When the lower extremities are used, the greater saphenous vein is often selected in pediatric patients because of its large size and consistent anatomy. It is not established whether femoral or jugular sites have fewer complications in neonates (25, 26).



FIG. 31.7. The jugular veins in relation to major anatomic landmarks.

- Catheter placement via jugular veins
 - Immobilize infant in position similar to that for percutaneous insertion of subclavian venous catheter.
 - If right side is to be catheterized, turn head to left and extend neck. Care must be taken not to extend the head too much, as this may result in occlusion of the neonatal vein.
 - Estimate length of catheter to be inserted by measuring from a point midway between the nipple and the midpoint of the clavicle to a point over the sternocleidomastoid muscle at the junction of the middle and lower third of the neck (Fig. 31.7).
 - Put on cap and mask.
 - Scrub as for major procedure and put on gown and gloves.
 - Prepare neck and scalp area or right chest wall with antiseptic solution such as iodophor and drape out the sterile field.
 - Make small, transverse incision (1 to 2 cm) through skin and platysma muscle low in the neck for the external jugular and higher up for the facial vein.
 - Free external jugular or facial vein by blunt dissection with curved mosquito hemostat. If internal jugular vein is used, sternocleidomastoid muscle must be split to locate vein.



FIG. 31.8. Catheterization of the external jugular vein; venotomy has been performed prior to inserting the catheter.

• Pass curved mosquito hemostat behind the vein, and place proximal and distal ligatures of 4-0 absorbable suture loosely around vein (Fig. 31.8). Be careful not to twist the vessels as the suture is advanced.



FIG. 31.9. Formation of a subcutaneous tunnel with a Vim-Silverman needle. A: Tunnel on the anterior chest wall. B: Alternative route under the scalp.

- Using a blunt tunneler, create a subcutaneous tract from neck to exit on the chest wall medial to the right nipple. In a baby girl, make sure that the tunnel is far from the breast bud (Figs. 31.9 and 31.10).
- Thread the end of the catheter through the opening in the tunneler, and guide the catheter gently through the subcutaneous tract.
- \circ $\;$ Fill the catheter system with heparinized flush solution.
- Cut the catheter length to the premeasured distance between the neck incision and a point midway between the center of the nipple line and the suprasternal notch.
- Perform transverse venotomy (Fig. 31.8).

For external jugular or facial vein (27)

- Tie cephalad venous ligature, and exert traction on both ligatures in opposite directions with aid of appropriately prepared assistant.
- Make short, transverse incision in anterior wall of vein, and enlarge gently by inserting and spreading tips of fine vascular forceps.

For internal jugular vein

- To avoid ligation of the vessel, use purse-string suture of 6-0 polypropylene, placed in vessel wall around point of catheter entrance.
- Make incision in vessel as for external jugular vein.
- \circ Bevel intravascular end of catheter (optional).
- Grasp catheter gently with blunt nontoothed tissue forceps, introduce catheter tip, and insert into the vein.
- Leave loop of catheter in neck wound to dampen effect of head movement (Fig. 31.11).
- Close wound with subcuticular 5-0 absorbable suture, taking care not to penetrate the catheter.
- Secure the catheter to the skin with at least one nylon suture to hold it until the cuff has created enough tissue ingrowth.
- Use selected method for fixation and dressing.



FIG. 31.10. Broviac catheter with transparent dressing.

- Proximal saphenous vein cutdown
 - Scrub and prepare as for major procedure.
 - Prepare as for cutdown on jugular vein.
 - Choose right or left groin area for insertion.
 - Prepare groin and abdomen on same side.
 - Make incision 1 cm long: 1 cm caudad and 1 cm lateral to pubic tubercle (Fig. 31.12).
 - Spread incision into subcutaneous tissues, using curved mosquito hemostat.
 - Incise superficial fascia.
 - Identify saphenous vein lying medial and inferior to its junction with femoral vein at foramen ovale (Fig. 31.12).
 - Move 0.5 to 1 cm distally before
 - Passing curved mosquito hemostat behind vein. This avoids inadvertent damage to femoral vein.
 - Placing two 4-0 absorbable suture ligatures loosely around vein
 - Create tunnel, using small hemostat or tunneling instrument, in subcutaneous plane laterally onto abdomen, just above or lateral to umbilicus or on lateral thigh.
 - \circ Flush catheter with heparinized saline and replace cap.



FIG. 31.11. Insertion of a catheter into the common facial vein. Incision is below the angle of the mandible at the level of the hyoid bone. The facial vein is ligated at the junction of the anterior and posterior tributaries. Inset: The catheter is looped in the neck wound to dampen the effect of head movement. Alternatively, a subcutaneous tunnel may be made with a catheter exit site on the anterior chest wall. (Reproduced from Zumbro GL Jr, Mullin MJ, Nelson TG. Catheter placement in infants needing total parenteral nutrition utilizing common facial vein. Arch Surg. 1971;102:71, with permission of American Medical Association.)



FIG. 31.12. Anatomic view of the site of incision for proximal saphenous vein cutdown with underlying femoral triangle.

• Pull catheter through tunnel into groin wound so that Dacron cuff is just within the skin incision.

Estimate length of catheter to be inserted so that tip will be in inferior vena cava at junction with right atrium.

- Cut catheter to appropriate length, and bevel intravascular end (optional).
- Dissect saphenous vein to junction with common femoral vein.
- Visualizing the junction prevents inadvertent direction of catheter into lower extremity.
- Apply traction to vein, using caudad suture. Lateral tension may also be applied by a scrubbed assistant, using fine nontoothed vascular forceps.
- Make transverse venotomy.
- o Dilate vein, if necessary, with blunt dilatator.
- Moisten catheter with saline to ease passage into vein.
- Maintain back-traction on caudad suture to control bleeding.
- Visualize catheter entering common femoral vein to ensure cephalad direction of catheter.
- Obtain radiograph to confirm position in inferior vena cava, once estimated length is inserted (radiographic contrast material may be required).
- Ligate vessel with caudad suture, and tie down cephalad suture without occluding catheter.
- Check for easy backflow of blood in catheter.
- Flush catheter with 2.5 to 3 mL of heparinized saline. If catheter is capped, while infant is transferred from operating room to intensive care unit, clamp catheter while plunger of heparin syringe is moving forward to ensure positive pressure in line to prevent backflow and clotting of blood.

- Close groin wound with subcuticular 5-0 absorbable suture, taking care not to penetrate catheter with needle.
- Secure the catheter to the skin with at least one nylon suture to hold it until the cuff has created enough tissue ingrowth.
- Cover with dressing of choice.

E. Sterile Dressing for Surgically Placed Central Venous Lines

Routine changing of central venous catheter dressings depends on the type of dressing. Transparent dressings should be changed at least every 7 days, and gauze dressings every 2 days. All dressings should be changed when damp, loose, or soiled (22).

Equipment

Strict sterile technique is used for all central line dressings.

- Antiseptic skin prep solution: Per institutional policy (e.g., 10% povidone–iodine or 0.5% chlorhexidine solution)
- Sterile gloves, mask, cap, and sterile gown (optional)
- Scissors (optional)
- Cotton-tipped applicator
- $4 \tilde{A} 4$ -in sterile gauze square
- Dressing of choice
 - Semipermeable transparent dressing
 - \circ Sterile 2- \tilde{A} 2-in gauze squares or presplit 2- \tilde{A} 2-cm gauze dressing
- Normal saline or sterile water
- Adhesive tape (if sterile tape not available, use fresh unused roll)

Precautions

- Procedure should be undertaken by trained personnel.
- Ensure that all personnel wear masks if within 6-ft radius of sterile area.
- Use strict aseptic technique.
- Remove dressing with care, to avoid cutting or dislodging catheter.
- If it is necessary to clamp the catheter, close the clamp on the catheter according to the manufacturer's directions. If the catheter does not have a clamp, use a rubber-shod clamp. Never place a clamp directly on the catheter.
- Never advance a dislodged catheter into the patient.
- Do not place adhesive tape on silicone tubing because this may occlude or damage the catheter.
- Do not routinely apply prophylactic topical antimicrobial or antiseptic ointment at the insertion site because of the potential for promoting fungal infections and antimicrobial resistance (22).

Technique

When a subcutaneous tunnel is used, occlusive dressing should be applied to both the cutdown site and the catheter exit site.

- Restrain patient appropriately, utilizing nonpharmacologic comfort measures.
- Put on head cover and mask.
- Scrub as for major procedure.
- Put on gown and gloves.
- Prepare sterile work area, using "no-touch†technique.
- Remove old dressing and discard.
- Inspect catheter site carefully (Table 31.4).
- Culture site if there is drainage or it appears inflamed.

- If area around catheter is contaminated with dried blood or drainage, clean with diluted hydrogen peroxide/sterile water solution (1:1).
- Remove gloves. Don sterile gloves.
- Cleanse area with antiseptic solution, starting at catheter site and working outward in circular motion for 2 to 4 cm. Repeat twice. Allow area to dry.
- Remove antiseptic with sterile water or saline gauze and allow to dry.



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FIG. 31.13. Occlusive dressing for a central venous line using presplit gauze. A: Placing split gauze over the skin entry site. B: Covering split gauze and the catheter with sterile gauze. Entire dressing is then covered with adhesive tape or clear dressing.

• Apply dressing of choice.

- Clear, adhesive, hypoallergenic, transparent dressing allows for continuous inspection of catheter insertion site (Fig 31.10).
 - If necessary, cut dressing to desired size.
 - Anchor dressing to skin above catheter skin entry site, so that the point of skin entry is at the center of the dressing.
 - Remove remainder of adhesive backing while applying dressing smoothly over site.
- Standard occlusive gauze dressing
 - Cut gauze halfway across, or use presplit gauze. Place around catheter as shown in Fig. 31.13.
 - Cover remainder of external catheter length (not hub) with sterile gauze.

- If sterile tape is not available, discard outer layer of tape on roll.
- Cover gauze with tape.
- Label dressing with initials and date.
- Secure intravenous tubing with tape to prevent tension on the center (a stress loop can decrease tension on the catheter).

F. Care of the Catheter When Not in Use for Continuous Infusion

Indications

To maintain patency and prevent clotting of the catheter when the line is used intermittently. Only largebore catheters (2.5 Fr or larger) may be kept patent by this technique. PICC lines that are 2 Fr or smaller tend to clot easily if continuous infusions are interrupted. Equipment

- 3 mL of heparin–saline solution (10 U/mL) in a 10-mL syringe (follow manufacturer's guidelines for syringe sizes)
- Alcohol wipes
- Catheter clamps (must have no teeth or be padded), or use clamp provided on catheter (Fig. 31.10)
- Clean gloves
- Intravenous injection cap (needleless is recommended)

Technique

- Converting to a heparin lock
 - Wash hands thoroughly.
 - Don clean gloves.
 - Prepare sterile work area.
 - Using aseptic technique, open sterile injection cap package and prefill injection cap with heparinized saline.
 - Clean the outside of the hub–intravenous tubing connection with an antiseptic such as alcohol wipes. Work outward in both directions. Allow to dry.
 - \circ $\,$ Clamp catheter with padded hemostat, or close catheter clamp.
 - Holding hub with alcohol swab, disconnect catheter hub from intravenous tubing.
 - Connect preflushed injection cap into hub of catheter (gently flushing during connecting can prevent air from entering catheter).
 - Release clamp and flush line with 1 to 3 mL of heparinized saline (depending on size of catheter).
 - Reclamp catheter while plunger of heparin syringe is depressed to prevent blood from backing into catheter (positive pressure).
 - Secure catheter and tape to chest or abdomen.
 - Flush catheter with heparinized solution every 6 to 12 hours (per institution policy).
- Flushing catheters

Equipment is same as for heparin lock.

- Wash hands thoroughly.
- Put on gloves and prepare sterile work area.
- Prepare intravenous injection cap with antiseptic solution. Allow to dry.
- If injection cap is part of a needleless system (recommended), connect flush syringe to cap. If the cap is not a needleless device, insert needle into intravenous catheter plug. Always use a 1-in needle. A longer needle can puncture the catheter.

- Unclamp catheter and slowly inject 1 to 2 mL of heparin solution (depending on catheter size). Reclamp catheter while injecting solution to prevent blood from flowing back into catheter. Positive-pressure injection caps are available to prevent backflow.
- Changing intravenous catheter injection cap: Most manufacturers recommend changing injection caps every 3 to 7 days, after blood product administration, or when they appear damaged (see specific manufacturer's instructions).

Catheter Removal

A. Indications

- Patient's condition no longer necessitates use.
- Occluded catheter
- Local infection/phlebitis
- Sepsis and/or positive blood cultures obtained through the catheter (catheter colonization). There are rare clinical circumstances when a catheter is left in place despite sepsis and antibiotic or antifungal therapy is administered through it in an attempt to clear the infection, but this may be associated with an increased risk of morbidity and mortality (28, 29).

B. Technique

Surgically implanted central venous catheters should be removed by a physician or other person specifically trained to remove cuffed and/or tunneled catheters.

- Remove dressing.
- Pull catheter from vessel slowly over 2 to 3 minutes.

Avoid excessive traction if catheter is tethered, because the catheter may snap (see Complications).

- Apply continuous pressure to the catheter insertion site for 5 to 10 minutes, until no bleeding is noted.
- Inspect catheter (without contaminating tip) to ensure that entire length has been removed.
- The cuff on the tunneled catheter should be dissected out under local anesthesia with IV sedation. If cuffs are retained, they may rarely cause more than a persistent small subcutaneous lump, although they can occasionally extrude through the skin.
- If desired, antibiotic ointment may be placed over site.
- Dress with small, self-adhesive bandage or gauze pad and inspect daily until healing occurs.

Complications of Central Venous Lines

- Damage to other vessels and organs during insertion
 - Possible during both percutaneous and surgical placement of central venous catheters
 - Complications include bleeding, pneumothorax, pneumomediastinum, hemothorax, arterial puncture, and brachial plexus injury (20, 30, 31).
- Phlebitis
 - Mechanical phlebitis may occur in the first 24 hours after line placement as a normal response of the body to the irritation of the catheter in the vein.
 - Management of mild phlebitis (mild erythema and/or edema): Apply moist, warm compress, and elevate extremity.
 - Remove the catheter if symptoms do not improve, if phlebitis is severe (streak formation, palpable venous cord, and/or purulent drainage), or if there are signs of a catheter-related infection.

- Catheter migration/malposition (2, 8, 9, 10, 11, 12, 13, 16) (Fig 31.14)
 - Can occur during insertion or from spontaneous migration at any point during the catheter dwell time. During insertion, the catheter can enter a side vein or reverse direction, causing it to loop or curl backward.
 - Sites of misplacement include the cardiac chambers, internal jugular vein, contralateral subclavian vein, ascending lumbar vein, and others, with consequences such as pericardial effusion or pleural effusion, cardiac arrhythmias, tissue extravasation/infiltration, and thrombosis.
 - The decision to remove the catheter or attempt to correct the position is based on the location of the tip. Although PICCs are intended to be placed in central veins, occasionally, the tip is in a noncentral location (e.g., in the subclavian vein). These noncentral PICCs may be used temporarily, provided the fluids administered through them are isotonic, but the care of the catheters must be as stringent as for centrally placed catheters (32).
 - The catheter should be pulled back a few centimeters if the tip is in the heart, as serious consequences such as cardiac arrhythmia, perforation, or pericardial effusion can occur.
 - Spontaneous correction of malpositioned lines has been demonstrated in some cases (33). If the tip of the catheter is looped into the internal jugular or in the contralateral brachiocephalic vein, the catheter may be used temporarily (using isotonic fluids that are suitable for peripheral venous cannulae) and re-evaluated radiologically in 24 hours. If the catheter has not moved spontaneously into the desired location, it should be removed.
- Infection (most common complication)
 - Catheter-related sepsis (CRS) rates range from 0% to 29% of lines placed and from 2 to 49 per 1,000 catheter days, with the smallest and most immature infants being at greatest risk (2, 34, 32, 36).
 - Strict protocols for central line care and a methodology of surveillance with a data feedback mechanism are recommended to decrease the rate of infection, which can vary significantly among neonatal intensive care units (4, 22, 34, 36).
 - Management of catheter-related sepsis: Remove central venous line if possible. Prompt removal of the line is recommended for Staphylococcus aureus, gram-negative, or Candida sepsis. Treatment with appropriate antibiotics without removal of the line may be attempted in infants with coagulase-negative Staphylococcus sepsis, but repeated positive cultures mandate removal of the line (28, 29).
- Catheter dysfunction
 - Obstruction of the catheter is characterized by increased pump pressures, or inability to infuse fluids or withdraw blood.
 - Dysfunction may be due to malposition, fibrin thrombosis, precipitates caused by minerals or drugs, or lipid deposits (36, 37).
 - Management
 - Check catheter position on chest radiograph.
 - If malposition is ruled out, review history of fluids and drugs administered through the catheter to determine probable cause of occlusion.
 - Remove the catheter if it is no longer medically critical.
 - Attempt clot dissolution only if maintenance of catheter is essential.
 - Equipment required: Face mask, sterile gloves and drape, prep solution, sterile threeway stopcock, a 10-mL syringe, and a 3-mL syringe filled with 0.2 to 0.5 mL of agent for clot dissolution.
 - Agents for clot dissolution (37)
 - Hydrochloric acid, 0.1 N, for calcium salt precipitates or drugs with pH < 7



FIG. 31.14. Various venous malpositions of subclavian venous catheters. A: Jugular. B: Looped in the right atrium with the tip in the superior vena cava. C: Looped in the superior vena cava. D: Looped in the innominate vein with the tip overlying the left scapula. E: Knotted in the left atrium.

- Sodium bicarbonate, 8.4%, 1 mEq/mL, for medications with pH > 7
- Ethanol, 70% concentration, for lipid deposits
- Recombinant tissue plasminogen activator, 0.5 to 1 mg/mL, for fibrin or blood clot (37, 38, 39)
- Recombinant urokinase, 2,000 to 5,000 IU/mL, for fibrin or blood clot (40)
- Technique
 - Use strict aseptic technique.
 - Remove IV tubing and cap to maintain sterility. After cleaning with prep, attach a three-way stopcock to catheter hub.

- Attach an empty 10-mL syringe to the side port of the three-way stopcock and a prefilled 3-mL syringe to the other port. Avoid use of 1-mL tuberculin syringe.
- Turn the stopcock off toward the prefilled syringe and open toward the empty syringe.
- Aspirate on the empty syringe, creating negative pressure in the occluded catheter.
- While maintaining the negative pressure, turn the stopcock off to the empty syringe and open to the prefilled syringe. The negative pressure in the catheter will automatically cause the medication in the prefilled syringe to flow into the catheter.
- Allow the medication to dwell in the catheter for 20 minutes to 1 hour.
- Aspirate after the dwell time to check for blood return, discard the aspirate, and flush the catheter with sterile normal saline. Resume catheter use.
- If the procedure is unsuccessful, it may be repeated once, or a different declotting agent may be tried.
- Do not use hydrochloric acid immediately before or after using sodium bicarbonate.
- Continuous infusions of low-dose recombinant tissue plasminogen activator or urokinase may be attempted if bolus doses do not clear the occlusion. Risk of hemorrhagic complications should be evaluated (41, 42).
- Thrombosis, thromboembolism
 - About 90% of venous thromboembolic events in neonates are associated with central venous catheters (43). They include:
 - Deep venous thrombosis
 - Superior vena cava syndrome
 - Intracardiac thrombus
 - Pulmonary embolism
 - Renal vein thrombosis
 - The complications of venous thrombosis include loss of venous access, potential danger of injury to affected organ or limb, thrombus propagation, embolization to other areas, and infection.
 - Management of thromboembolism in neonates is controversial. The severity of thrombosis and the potential risk to organs or limbs dictate the degree of intervention required, including the use of thrombolytic/anticoagulant therapy or surgical intervention (44).
- Extravascular collection of fluid
 - Pericardial effusion with or without cardiac tamponade (Fig. 37.3) (2, 8, 9, 10, 11, 12, 13). This serious complication presents as sudden cardiac collapse or unexplained cardiorespiratory instability. The cardiothoracic ratio is increased, and pulsus paradoxus may be noted (Fig. 37.1). Immediate pericardiocentesis may be life-saving (Chapter 37).
 - Pleural effusion
 - $\circ \quad \mbox{Mediastinal extravasation}$
 - Hemothorax
 - \circ Chylothorax
 - Ascites
- Catheter breakage
 - Catheters may be severed by the introducer needle during insertion of a PICC, snap because of excessive tension on the external portion of the catheter, or rupture because of excessive pressure. Other common causes include external clamps, kinking of the catheter, constricting sutures, and poorly secured catheters. The intravascular portion of the broken catheter is at risk for embolization (45).

- In the event of catheter breakage, immediately grasp and secure the extravascular portion of the broken catheter to prevent migration.
- If the catheter is not visible outside the baby, apply pressure over the venous tract above the insertion site to prevent the catheter from advancing. Immobilize the infant, and obtain a radiograph immediately to localize the catheter.
- Surgical and/or cardiothoracic intervention may be required if the catheter is not visible externally.
- Damaged or broken catheters must be removed and replaced. Repaired catheters and catheter replacement over a guidewire place the patient at risk for infection or embolization. If no other options exist owing to limited venous access, the catheter can sometimes be repaired, utilizing meticulous aseptic technique. Repaired PICCs should be considered temporary, and a new catheter should be placed as soon as possible. Some manufacturers offer repair kits and instructions. A butterfly or blunt needle may be used in an emergency (46) (Fig. 31.15).
- Tethered catheter
 - Difficulty in removing catheter may be due to the formation of a fibrin sheath or secondary to sepsis.
 - Management
 - Place warm compresses on skin along the vein.
 - Use gradual, gentle traction on the catheter.
 - Thrombolytic therapy (47)
 - Surgical removal through a peripheral incision



FIG. 31.15. Emergency catheter repair using butterfly needle (46). (From Neonatal Network, Santa Rosa, CA, USA, with permission.)

References

 Ainsworth SB, Clerihew L, McGuire W. Percutaneous central venous catheters versus peripheral cannulae for delivery of parenteral nutrition in newborns. Cochrane Database Syst Rev. 2004(2):CD004219.
 Cartwright DW. Central venous lines in neonates: a study of 2186 catheters. Arch Dis Child Fetal Neonatal Ed. 2004;89:504–508.

3. Mahieu LM, De Dooy JJ, Lenaerts AE, et al. Catheter manipulations and the risk of catheter-associated bloodstream infection in neonatal intensive care unit patients. J Hosp Inf. 2001;48:20–26.

4. Schelonka RL, Scruggs S, Nichols K, et al. Sustained reductions in neonatal nosocomial infection rates following a comprehensive infection control intervention. J Perinatol. 2006;26:141–143.

5. Rodrigues AF, van Mourik ID, Sharif K, et al. Management of end-stage central venous access in children referred for possible small bowel transplantation. J Pediatr Gastroenterol Nutr. 2006;42:427–433.
6. Azizkhan RG, Taylor LA, Jacques PF, et al. Percutaneous translumbar and transhepatic inferior vena caval catheters for prolonged vascular access in children. J Pediatr Surg. 1992;27: 165–169.

7. Fletcher SJ, Bodenham AR. Safe placement of central venous catheters: where should the tip of the catheter lie? Br J Anaesth. 2000;85:188–191.

8. Nowlen TT, Rosenthal GL, Johnson GL, et al. Pericardial effusion and tamponade in infants with central catheters. Pediatrics. 2002;110:137–142.

9. Beardsall K, White DK, Pinto EM, et al. Pericardial effusion and cardiac tamponade as complications of neonatal long lines: are they really a problem? Arch Dis Child Fetal Neonatal Ed. 2003;88:292–295.

10. Jouvencel P, Tourneux P, Perez T, et al. Central catheters and pericardial effusion: results of a multicentric retrospective study. Arch Pediatr. 2005;12:1456–1461.

11. Darling JC, Newell SJ, Mohamdee O, et al. Central venous catheter tip in the right atrium: a risk factor for neonatal cardiac tamponade. J Perinatol. 2001;21:461–464.

12. Chock VY. Therapeutic techniques: peripherally inserted central catheters in neonates. NeoReviews. 2004;5:61–62.

13. Pezzati M, Filippi L, Chiti G, et al. Central venous catheters and cardiac tamponade in preterm infants. Intensive Care Med. 2004;30:2253–2256.

14. Odd DE, Battin MR, Kuschel CA. Variation in identifying neonatal percutaneous central venous line position. J Paediatr Child Health. 2004;40:540–543.

15. Webster NJ, Page B, Kuschel CA, Battin MR. Digital imaging does not improve localization of percutaneously inserted central lines in neonates. J Paediatr Child Health. 2005;41:256–259.

16. Coit AK, Kamitsuka MD. Peripherally inserted central catheter using the saphenous vein: importance of two-view radiographs to determine the tip location. J Perinatol. 2005;25:674–676.

17. Odd DE, Page B, Battin MR, Harding JE. Does radio-opaque contrast improve radiographic localisation of percutaneous central venous lines? Arch Dis Child Fetal Neonatal Ed. 2004;89: 41–43.

18. Reece A, Ubhi T, Craig AR, Newell SJ. Positioning long lines: contrast versus plain radiography. Arch Dis Child Fetal Neonatal Ed. 2001;84:129–130.

19. Groves AM, Kuschel CA, Battin MR. Neonatal long lines: localization with colour Doppler ultrasonography. Arch Dis Child Fetal Neonatal Ed. 2005;90:F5.

20. Casado-Flores J, Barja J, Martino R, et al. Complications of central venous catheterization in critically ill children. Pediatr Crit Care Med. 2001;2:57–62.

21. Primhak RH, Gathercole N, Reiter H. Pressures used to flush venous catheter. Arch Dis Child Fetal Neonatal Ed. 1998;78:F234.

22. U.S. Department of Health and Human Services. Centers for Disease Control and Prevention. Guideline for prevention of intravascular device-related infection. Morbidity and Mortality Report. 2002:51(RR-10).
23. Shah P, Shah V. Continuous heparin infusion to prevent thrombosis and catheter occlusion in neonates with peripherally placed percutaneous central venous catheters. Cochrane Database Syst Rev.

2005;20(3):CD002772.

24. Wong EC, Schreiber S, Criss V, et al. Feasibility of red blood cell transfusions through small bore central venous catheters in neonates. Pediatr Res. 2001;49:322A.

25. Vegunta RK, Loethen P, Wallace LJ, et al. Differences in the outcome of surgically placed long-term central venous catheters in neonates: neck vs groin placement. J Pediatr Surg. 2005;40(1): 47–51.

26. Murai DT. Are femoral Broviac catheters effective and safe? A prospective comparison of femoral and jugular venous Broviac catheters in newborn infants. Chest. 2002;121:1527–1530.

27. Zumbro GL Jr, Mullin MJ, Nelson TG. Catheter placement in infants needing total parenteral nutrition utilizing common facial vein. Arch Surg. 1971;102:71.

28. Benjamin DK, Miller W, Garges H, et al. Bacteremia, central catheters, and neonates: when to pull the line. Pediatrics. 2001;107(6):1272–1276.

29. Karlowicz MG, Furigay PJ, Croitoru DP, Buescher ES. Central venous catheter removal versus in situ treatment in neonates with coagulase-negative staphylococcal bacteremia. Pediatr Infect Dis J. 2002;21(1):22–27.

30. Pandit PB, Pandit FA, Govan J, et al. Complications associated with surgically placed central venous catheters in low birth weight neonates. J Perinatol. 1999;19:106–109.

31. Pleasure JR, Shashikumar VL. Phrenic nerve damage in the tiny infant during vein cannulation for parenteral nutrition. Am J Perinatol. 1990;7(2):136.

32. Thiagarajan RR, Bratton SL, Gettmann T, Ramamoorthy C. Efficacy of peripherally inserted central venous catheters placed in noncentral veins. Arch Pediatr Adolesc Med. 1998;152(5): 436–439.

33. Rastogi S, Bhutada A, Sahni R, et al. Spontaneous correction of the malpositioned percutaneous central venous line in infants. Pediatr Radiol. 1998;28:694–696.

34. Aly H, Herson V, Duncan A, et al. Is bloodstream infection preventable among premature infants? A tale of two cities. Pediatrics. 2005;115:1513–1518.

35. Chien L, Macnab Y, Aziz K, et al. Variations in central venous catheter-related infection risks among Canadian neonatal intensive care units. Pediatr Infect Dis J. 2002;21(6):505–511.

36. Petit J. Assessment of infant with peripherally inserted central catheters: Part 1. Detecting the most frequently occurring complications. Adv Neonatal Care. 2002;2(6):304–315.

37. Kerner JA, Garcia-Careaga MG, Fisher AA, Poole RL. Treatment of catheter occlusion in pediatric patients. J Parenter Enteral Nutr. 2006;30(1 suppl):S73–S81.

38. Lee EK. Alteplase use for prevention of central line occlusion in a preterm infant. Ann Pharmacother. 2002;36(2):272–274.

39. Jacobs BR, Haygood M, Hingl J. Recombinant tissue plasminogen activator in the treatment of central venous catheter occlusion in children. J Pediatr. 2001;139(4):593–596.

40. Svoboda P, Barton RP, Barbarash OL, et al. Recombinant urokinase is safe and effective in restoring patency to occluded central venous access devices: a multiple-center, international trial. Crit Care Med. 2004;32(10):1990–1996.

41. Hooke C. Recombinant tissue plasminogen activator for central venous access device occlusion. J Pediatr Oncol Nurs. 2000;17: 174–178.

42. Bagnall HA, Gomperts E, Atkinson JB. Continuous infusion of low dose urokinase in the treatment of central venous catheter thrombosis in infants and children. Pediatrics. 1989;83:963–966.

43. Schmidt B, Andrew M. Neonatal thrombosis: report of a prospective Canadian and international registry. Pediatrics. 1995;96(5):939–943.

44. Ramasethu J. Management of vascular thrombosis and spasm in the newborn. Neoreviews. 2005;6:e1–e14.

45. Chow LML, Friedman JN, MacArthur C, et al. Peripherally inserted central catheter (PICC) fracture and embolization in the pediatric population. J Pediatr. 2003;142:141–144.

46. Evans M, Lentsch D. Percutaneously inserted polyurethane central catheters in the NICU: one unit's experience. Neonatal Network. 1999;18:37–46.

47. Nguyen ST, Lund CH, Durand DJ. Thrombolytic therapy for adhesions of percutaneous central venous catheters to vein intima associated with Malassezia furfur infection. J Perinatol. 2001;21:331–333.

32-Extracorporeal Membrane Oxygenation Cannulation and Decannulation

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Extracorporeal membrane oxygenation (ECMO) is defined as the use of a modified heart–lung machine combined with a membrane oxygenator to provide cardiopulmonary support for patients with reversible pulmonary and/or cardiac insufficiency in whom maximal conventional therapies have failed (1,2,3). After decades of laboratory and clinical research, ECMO is now well accepted as a standard of treatment for neonatal respiratory failure refractory to conventional techniques of pulmonary support (4,5,6,7). Most causes of neonatal respiratory failure are self-limited, and ECMO allows time for the lung to recover from the underlying disease process and for reversal of pulmonary hypertension, which frequently accompanies respiratory failure in the newborn.

Venoarterial Extracorporeal Membrane Oxygenationâ€"Cannulation

A. Indications

Placement of carotid arterial and internal jugular venous catheters for use in venoarterial ECMO.Venoarterial ECMO should be used in patients with significant cardiovascular instability.B. Relative Contraindications for Extracorporeal Membrane Oxygenation in the Neonatal Period (5,7)

- Gestational age <34 weeks
- Birthweight <2,000 g
- Uncontrolled coagulopathy or bleeding disorders
- Congenital heart disease without lung disease. Exception: Postoperative cardiac patients, a topic that will not be covered in this chapter
- Irreversible lung pathology
- Intracranial hemorrhage >Grade I to II
- Major lethal congenital anomaly
- Duration of maximum ventilatory support, >10 to 14 days
- Significant positive response to ventilator management and/or inhaled nitric oxide (iNO)

C. Precautions

- Ensure that the patient is paralyzed before placing the venous catheter, to prevent air embolus.
- Recognize that:
 - Internal jugular lines placed for intravenous access prior to ECMO may cause clot formation, resulting in the need for thrombectomy before placement of the venous ECMO catheter.
 - Excessive manipulation of the internal jugular vein may cause spasm and inability to place a catheter of appropriate gauge.
 - A lacerated vessel may result in the need for a sternotomy for vessel retrieval.

Appropriate instruments should be on the bedside tray or cart. A backup unit of blood should be available in the blood bank.

• Blood loss sufficient to produce hypotension can occur during a difficult cannulation.

Emergency blood should be available at the bedside (10 to 20 mL/kg).

- The vagus nerve is located next to the neck vessels and may be injured or manipulated during isolation of the vessels. Manipulation can cause bradycardia or other arrhythmias.
- Vital signs and pulse oximetry values must be monitored at all times because clinical observation of the infant is prevented by the surgical drapes.
- If the patient has been hand bag–ventilated for stabilization, do not place the Ambu bag on the bedside when surgical drapes are placed. The bag may entrap oxygen, which can result in a fire when electrocautery is used.

D. Personnel, Equipment, and Medications (8)

Personnel

- Surgical team
 - A senior surgeon (pediatric, cardiovascular, or thoracic) with assistant
 - A surgical scrub nurse and a circulating nurse
- Medical team
 - A physician trained in management of ECMO patients and cannulation techniques, who will administer anesthetic agents and manage the infant medically during the procedure
 - A bedside intensive care (neonatal or pediatric intensive care unit) nurse, who will monitor vital signs, record events, and draw up medications as needed by the ECMO physician
 - A respiratory therapist, who will change ventilator settings as necessary
- Circuit specialists
 - A cardiovascular perfusionist, nurse, or respiratory therapist specially trained in this procedure, who will prime the pump
 - A bedside ECMO specialist (nurse, respiratory therapist, or cardiovascular perfusionist with special training in ECMO management), who will manage the ECMO system after the patient is on ECMO

Equipment (Fig. 32.1) Sterile

- Arterial and venous catheters (9)
 - Arterial
 - The size of the arterial catheter determines the resistance of the ECMO circuit, because it is the part of the ECMO circuit with the smallest internal diameter and thus the highest resistance.
 - This catheter should be as short as possible, with a thin wall and a large internal diameter (resistance is related directly to the length of the catheter and inversely to the diameter). An example of a suitable catheter is the Bio-Medicus Extracorporeal Circulation Cannula, 8 to 10 French (Fr) (Bio-Medicus, Minneapolis, MN, USA).
 - Venous
 - The venous catheter should have as large an internal diameter as possible to allow maximal blood flow (the patient's oxygenation is related directly to the rate of blood flow).
 - Should be thin walled with a large internal diameter. An example of a suitable catheter is the Bio-Medicus Extracorporeal Circulation Cannula, 8 to 14 Fr. (Bio-Medicus, Minneapolis, MN, USA).



FIG. 32.1. Schematic diagram of venoarterial extracorporeal membrane oxygenation circuit, showing the drainage from the right atrium into the bladder of the circuit, with flow through the membrane lung, heat exchanger, and return flow to the arch of the aorta via the carotid artery catheter. (From Polin RA, Fox WC, eds. Fetal and Neonatal Physiology, Vol. 1. Philadelphia: WB Saunders; 1992:933, with permission.)

- Surgical instruments required are listed in Tables 32.1 and 32.2.
- Sterile gowns and gloves
- Sterile saline for injection
- Syringes (1 to 20 mL) and needles (19 to 26 gauge)
- Povidone–iodine solution
- Povidone iodine ointment
- Semipermeable transparent membrane-type dressing
- Absorbable gelatin sponge, for example, Gelfoam (Upjohn, Kalamazoo, MI, USA)
- Surgical lubricant, bacteriostatic

Nonsterile

- Surgical head covers and mask
- Pulse oximeter
- Surgical head light
- Electrocautery
- Wall suction
- Shoulder roll, for example, a small blanket, to place under infant's shoulders
- Tubing clamps

Medications

- A long-acting paralyzing agent, for example, pancuronium bromide (0.1 mg/kg)
- Fentanyl citrate (10 to 20 mcg/kg)

- Sodium heparin (75 to 150 U/kg)
- Topical thrombin/Gelfoam
- Xylocaine, 0.25%, with epinephrine
- Xylocaine, 1%, plain (without epinephrine)
- Cryoprecipitate, thawed, or commercially available fibrin sealant (optional)

TABLE 32.1 Surgical Instruments for ECMO Cannulation

Number	Item
Place in a 1	2 - 18-in mayo tray with a Huck towel on the bottom of the tray.
2	Custard cup (place on inside of other cup with a 3 - 4-in sponge)
1	Medicine cup (place inside of custard cup with a 3 - 4-in sponge)
2	Straight bulldog clamps
1	Sauer eye retractor
1	Alm retractor
1	Mastoid Jansen retractor
2	Vein retractors
2	Octagonal forceps
2	7-in Gerald forceps
2	6-in DeBakey forceps
1	Adson forceps, plain
2	Adson forceps with teeth
2	No. 3 knife handles
1	Castroviejo needleholder
2	Right-angle retractors
2	Chops retractors
1	Set of Garrett dilators, nine pieces (sizes 1.0, 1.5, 2.0, 2.5, 3.0, 3.5, 4.0, 4.5, 5.0)
String the fo	ollowing instruments from left to right on two 9-in sponge sticks or instrument stringer. Then
place on top	o of a rolled Huck towel.
4	9-in sponge stick
1	Tonsil clamp (bleeder)
1	612-in crile
1	534-in crile
1	Baby right-angle clamp
4	Straight mosquitoes
6	Curved mosquitoes
3	Fine curved mosquitoes
2	Tubing clamp with guard
1	Ryder needleholder
1	Webster needleholder
1	Straight mayo scissor
1	534-in Metzenbaum scissor
1	Curved Steven scissor
1	Straight Iris scissor
4	Small towel clips (nonpenetrating)
1	Baby Satinsky clamp
1	Curved bulldog clamp
1	Straight bulldog clamp
1	Disposable ECMO tray (Table 32.2)
ECMO, ext	racorporeal membrane oxygenation. For information on suture material, see Appendix B2.

Nun	nberItem
2	1-mL syringe
1	20-mL syringe
1	6-mL syringe
1	3-mL syringe
1	Needle adapter
3	Single-cavity tray
2	Gauze packages
1	Betadine ointment
1	Surgical blade no. 15 carbon
2	Semipermeable transparent dressings
1	Handle, suction frazier, 8 Fr
1	Xylocaine insert
1	Mini yellow vessel loops
1	Hand-control cautery
1	Suture, 4-0 Vicryl
1	Suture, 2-0 silk
1	Suture, 6-0 Prolene
4	Forceps, sponge
1	25-gauge needle
1	NaCl, 5-mL amp
1	3-g foil package of Surgilube
1	Surgical blade, no. 11 carbon
2	Steri-Drapes
2	Connectors, straight 14 — 14 in
1	Xylocaine 1%
1	Suction tubing, 316 in — 10 ft

TABLE 32.2 Contents of Disposable Extracorporeal Membrane Oxygenation Tray

E. Technique Preparation for Cannulation

Package sterile towels (14)

1

- Place infant with head to foot of warmer.
- Anesthetize the patient with fentanyl (10 to 20 mcg/kg).
- Paralyze the patient with pancuronium (0.1 mg/kg).
- Hyperextend the patient's neck with a shoulder roll, and turn the head to the left (Fig. 32.2). Make sure that the Bovie ground pad is placed at this time.

Observe closely for hypotension.

- Monitor vital signs and give additional fentanyl and/or pancuronium as needed (see Chapter 5).
- Clean a wide area of the right neck, chest, and ear with Betadine solution.
- Drape the infant and entire bed with sterile towels.
- Use Steri-Drapes (3M Health Care, St. Paul, MN, USA) to secure the towels to the skin.



FIG. 32.2. Infant positioned for cannulation with shoulder roll present and head extended to the left. Position of neck incision is indicated.

- At the point of incision, infiltrate the skin with Xylocaine (AstraZeneca, Wayne, PA, USA) (0.25%, with epinephrine) (Fig. 32.2).
- Wait at least 3 minutes for anesthesia to be effective.
- Make a 1- to 2-cm vertical incision over the right sternocleidomastoid muscle, starting approximately 1 cm above the right clavicular head, using the electrocautery set on cutting current (Fig. 32.3).
- Continue to use the electrocautery to cut through the subcutaneous tissue.



FIG. 32.3. Landmarks over the sternocleidomastoid muscle for making the incision with electrocautery.

- Coagulate all visible bleeding sites.
- Spread the fibers of the sternocleidomastoid muscle apart with a hemostat and retract using hemostats clamped onto the muscle (Fig. 32.4).
- Open the carotid sheath, taking care to avoid the vagus nerve.

• Irrigate both the common carotid artery and internal jugular vein with 1% plain Xylocaine to vasodilate the vessels.



FIG. 32.4. Split sternocleidomastoid and open carotid sheath.

- Encircle the artery with silicone loop, and proximal and distal 2-0 silk ties held with clamps but not tied. Avoid "sawing†the ties on the artery.
- Avoid excessive handling of the internal jugular vein. Some isolate the vein after cannulation of the carotid artery to avoid spasm.
- Estimate the length of the cannula to be inserted.
 - \circ $\;$ Identify the sternal notch and the xiphoid process.
 - The arterial catheter is inserted approximately one third of the distance between the sternal notch and the xiphoid process. This is typically between 3 and 4 cm.
 - The venous catheter is inserted approximately one half the distance between the sternal notch and the xiphoid process. This is typically between 7 and 7.5 cm.
 - Mark these distances on the catheters with a 2-0 tie, or note the distance if the cannula is marked.
 - Heparinize the patient with a bolus of 75 to 150 U/kg of heparin, depending on the estimated risk of bleeding, and wait 60 to 90 seconds before proceeding with cannulation.



FIG. 32.5. A: Carotid artery isolated with vessel clamp in place and with arteriotomy site showing the placement of the 6-0 Prolene traction sutures. B (inset): Magnified view of (A).

Arterial cannulation

• Tie the distal ligature on the carotid artery, and place a bulldog clamp on the proximal portion of the artery.

Allow blood to dilate the artery before placing the bulldog clamp.

• Make an arteriotomy using a no. 11 scalpel blade, and place two traction sutures of 6-0 Prolene (Ethicon, Somerville, NJ, USA) on the proximal side of the arteriotomy (Fig. 32.5).

Always use traction sutures, to prevent intimal tears.

- If desired, lubricate Garrett dilators with sterile surgical lubricant and dilate the artery to the approximate size of the catheter.
- Place a sterile tubing clamp on the catheter. Lubricate the catheter and insert the catheter into the vessel as the bulldog clamp is removed.
- Secure the catheter with a 2-0 silk ligature tied over a 0.5- to 1-cm vessel loop ("bootieâ€) (Fig. 32.6).
- Place a second 2-0 silk ligature. Tie the distal tie around the catheter, and then tie the distal and proximal ties together. Some surgeons place two ties proximally and one distally for added security.
- Allow blood to back up into the catheter to remove air.

Venous cannulation

• Dissect the vein free and isolate with two 2-0 silk ties.

Do not apply traction to the vein with the ties, to avoid spasm.

- Place a bulldog clamp on the proximal end of the vein, allowing blood to distend it. Then tie the distal end of the vein with the 2-0 silk ligature.
- Make a venotomy with a no. 11 scalpel blade, and place two stay sutures of 6-0 Prolene as traction sutures, as for arterial cannulation.
- Lubricate the venous catheter, place a sterile tubing clamp on the catheter, and dilate the venotomy.

• Insert the catheter as an assistant places traction on the proximal tie, and apply pressure over the liver to increase the backflow of blood out of the catheter (to decrease the risk of an air embolus).

There will be a slight impedance to catheter advancement at the thoracic inletâ€"pushing against resistance will tear the vein. Use gentle downward and posterior pressure.



FIG. 32.6. A: Securing the catheter with proximal and distal ties onto a $\hat{a} \in \hat{B}$ (inset): Magnified view of (A).

- Secure, as for the artery, and back blood into the catheter by pressing gently on the liver.
- If desired, pack the wound with absorbable gelatin sponge soaked in topical thrombin or commercially available topical fibrin sealant, to assist in hemostasis.

Cryoprecipitate and topical thrombin can be used to form a fibrin clot if dropped onto the field from separate syringes in a one-to-one concentration. Note: If they are mixed together in one syringe, they will form a solid clot in the syringe. A similar product is now commercially available as Tisseel-HV Fibrin Sealant (Baxter Hyland Division, Glendale, CA, USA).

• Confirm catheter placement by chest radiography and/or cardiac echocardiography, if the patient is stable (Fig. 32.7) (10). If the patient is unstable, he or she can be placed on ECMO and the radiograph taken when adequate oxygenation is achieved but prior to closing the surgical wound.



FIG. 32.7. Radiograph at cannulation, showing proper placement of the arterial and venous catheters. Note the radio-opaque dot indicating the end of the Bio-Medicus venous extracorporeal membrane oxygenation catheter (arrow).

Venovenous Extracorporeal Membrane Oxygenation Cannulation

More than 60% of neonatal ECMO patients reported in the ELSO registry have received treatment with venoarterial bypass (11). In neonates with respiratory failure, venoarterial ECMO is gradually being replaced by a venovenous (VV) technique, which uses a single double-lumen catheter (Fig. 32.8). The catheter is placed in the right atrium, where blood is drained and reinfused into the same chamber, thus requiring cannulation of only the right jugular vein, thus sparing the carotid artery. Other advantages of VV ECMO include maintenance of normal pulsatile blood flow, and the theoretical advantage that particles entering the ECMO circuit enter by way of the pulmonary rather than the systemic circulation. The design of the original VV catheter resulted in significant recirculation, limiting its use when ECMO flows >350 mL/min were required. Research by Rais-Bahrami et al. resulted in development of a new catheter design that significantly lowers the degree of recirculation (12). The double-lumen catheter should be placed within the right atrium, directing the oxygenated blood from the return lumen through the tricuspid valve to minimize recirculation. This catheter design in 12-, 15-, and 18-Fr sizes allows the use of VV ECMO in a greater number of infants (13).



FIG. 32.8. Schematic of the venovenous extracorporeal membrane oxygenation catheter placed in the midright atrium. (From Short BL. CNMC ECMO Training Manual. 2005, with permission).

A. Double-Lumen Venovenous Catheters

- Kendall 14-Fr catheter (Kendall Health Care Products, Mansfield, MA, USA)
- OriGen 12-, 15-, and 18-Fr catheters (OriGen Biomedical, Austin, TX, USA)
- B. Advantages of Venovenous Bypass
 - Provides excellent pulmonary support
 - Avoids carotid artery ligation
 - Oxygenated blood enters pulmonary circulation.
 - Particles coming from the ECMO circuit enter the venous circulation instead of the arterial circulation.
- C. Disadvantages of Venovenous Bypass
 - Lack of cardiac support
 - ECMO support is dependent on the patient's cardiac function.
 - Catheter position and rotation are extremely critical.
 - Amount of recirculation

D. Cannulation Technique

The cannulation technique for venovenous ECMO is essentially the same procedure as venous cannulation for venoarterial ECMO, with the following exceptions:

• Both internal jugular vein and carotid arteries are identified and dissected free, although the internal jugular vein is the only vessel cannulated with the double-lumen venovenous catheter. A silastic loop may be tied loosely around the artery to facilitate potential conversion to venoarterial flow.

Both vessels are isolated in case a rapid conversion to venoarterial bypass becomes necessary.

• The catheter is advanced in a direction with the arterial side upward and anterior to the venous limb of the double-lumen cannula.

Caution: Avoid bending the catheter or creating a "crimp†in the catheter.

Correct positioning of the catheter helps direct the oxygenated blood return toward the tricuspid valve, thus minimizing the recirculation of the oxygenated blood back to the ECMO circuit.

• The proximal end of the internal jugular vein is also cannulated for cephalad drainage, that is, a jugular bulb catheter. This catheter is connected to the venous tubing of the ECMO circuit via a Luer connector. For this, we use a custom-made Carmeda heparin-coated Bio-Medicus venous catheter, made specifically for use as a cephalad catheter.

This allows additional venous drainage to the ECMO circuit, prevents venous congestion, and also allows for cephalic venous saturation measurement.

• If using a jugular bulb catheter to measure cerebral saturations, care should be used when entering the circuit; air will draw into the venous side of the circuit rapidly if a stopcock is loose or is left open.

E. Placing Patient on the Extracorporeal Membrane Oxygenation Circuit

The circuit has been previously primed with packed cells/albumin. The priming procedure and the surgical placement of the ECMO catheters should be timed so that the two are completed at the same time. Priming of the circuit is beyond the scope of this chapter.

- Fill catheters with sterile saline. Connect them to the ECMO circuit by inserting the ¼X¼-in connectors into the tubing as the assistant drips sterile saline into the ends of the circuit tubing and the catheter, to ensure that all residual air is eliminated prior to connection.
 - Do not squeeze the tubing while attaching; air will enter when the tubing is released.
 - If air is seen in the tubing, the catheters must be disconnected from the circuit. Prior to reconnection, air is removed, and the catheters are reconnected as described in E.1.
- Remove all sterile tubing clamps from the catheters, and have a nonsterile assistant hold the catheters. Nonsterile tubing clamps remain in place on the arterial and venous sides of the circuit at this juncture.
- Place the patient on ECMO by removing the arterial clamp, placing a clamp on the bridge (Fig. 32.9a), and removing the venous clamp. This will remove all nonsterile clamps from the circuit.
- Increase ECMO flow in 50-mL increments over 20 to 30 minutes, until adequate oxygenation is achieved (usually at 120 mL/kg/min).

Transfusion may be needed if hypotension occurs at this stage.

• Decrease the ventilator settings and oxygen concentration gradually as the ECMO flows are increased.

Typical resting ventilator settings for venoarterial ECMO are at a rate of 10 to 15 breaths/min, a peak pressure limit of 15 to 20 cm H_2O , and F_iO_2 of 0.21 to 0.30. For venovenous ECMO, it is recommended to keep ventilator settings at a rate of 20 to 30 breaths/min, a peak inspiratory pressure of 20 to 25 cm H_2O , and F_iO_2 of 0.30 to 0.50.

F. Closure of the Neck Wound

• Obtain radiographic confirmation of appropriate catheter position and achievement of an adequate flow rate through the ECMO circuit prior to closure of the neck wound.

- Cut and remove traction sutures.
- Approximate the skin with a running 4-0 Vicryl (Ethicon) suture on an atraumatic needle.
- Tie the Vicryl suture, and use the tails of the suture to secure each catheter.
- Tie catheters together with another silk tie.
- Anesthetize the area behind the ear with 0.25% Xylocaine with epinephrine.
- Use 2-0 silk suture on a noncutting needle to place a stitch behind the ear and tie around the catheter to secure in place. Place a separate stitch for each catheter.
- Tie catheters together, dress the incision with povidone–iodine ointment, and cover the area with semipermeable membrane dressing.
- Tape the circuit tubing securely to the bedside to reduce traction on the catheters.

G. Complications

- Torn vessels, more commonly the vein
 - This risk is decreased if 6-0 Prolene stay sutures are always used.
 - Do not attempt to use too large a catheter.
- Aortic dissection associated with arterial cannulation (14)
- Blood loss, particularly during the venous cannulation, when side holes in the catheter are outside the vein
- Venous spasm, resulting in inability to place a large enough venous catheter to meet the required ECMO flow to support the patient adequately

The rate of blood flow is impeded by the small gauge of the catheter, requiring that a second venous catheter be placed in the femoral vein. The two catheters must be Y-connected together into the ECMO circuit.

- Arrhythmias and/or bradycardia can occur, owing to stimulation of the vagus nerve
- Hypotension, due to an increase in the intravascular space when the patient is connected to the ECMO circuit
- Conversion to venoarterial from venovenous ECMO. This will occur if:
 - The patient remains hypoxic despite adequate ECMO flow.
 - \circ $\;$ The patient remains hypotensive despite vasopressor support.
 - Cerebral venous saturations remain persistently <60% after adequate flows and ventilator management have been undertaken.

Converting from venovenous to venoarterial ECMO requires cannulation of carotid artery with a Biomedicus arterial catheter, and the double-lumen venovenous catheter must be $\hat{a} \in \mathfrak{C}Y'd\hat{a}$ in together to make a double-lumen venous drainage catheter (Fig. 32.9).

Extracorporeal Membrane Oxygenationâ€"Decannulation



FIG. 32.9. Schematic view of converting from venoarterial (A) to venovenous (B) ECMO. The doublelumen venovenous catheter is $\hat{a} \in \mathfrak{C}Y'd\hat{a}$ together to make a double-lumen venous drainage catheter.

A. Indications

- Removal from ECMO after lung recovery
- Removal from ECMO because of a complication such as uncontrolled bleeding or failure of lung recovery

B. Contraindications

All intensive support is being withdrawn, and permission for autopsy is obtained. It is usually optimal to remove the catheters during the autopsy.

C. Precautions

- The patient must be paralyzed during the removal of the venous catheter to avoid an air embolus.
- The vessels are fragile and may tear. A backup unit of blood should be available at the bedside.
- Delay removing catheter for 12 to 24 hours in cases in which there is a high risk of reoccurrence of pulmonary hypertensionâ€" for example, severe congenital diaphragmatic hernia.

D. Personnel, Equipment, and Medications

Personnel

Same as for cannulation, with the exception of the primer, which is not required

Equipment

Sterile

- Surgical tray with towels and suture as for cannulation
- Semipermeable transparent dressing
- Povidone–iodine ointment
- Syringes (1 to 20 mL) and needles (18 to 26 gauge)
- Unit of blood
- Absorbable gelatin sponge

Nonsterile Same as for cannulation

Medications

- Fentanyl (10 to 20 mcg/kg)
- Vecuronium bromide (0.2 mg/kg)

A short-acting paralyzing agent is preferred because of the relatively short duration of the procedure. This allows the infant to breathe spontaneously as soon as possible after decannulation, which facilitates rapid weaning from ventilator support.

- Xylocaine, 0.25%, with epinephrine
- Topical thrombin
- Protamine sulfate (1 mg only)

E. Technique

Postdecannulation vessel reconstruction is beyond the scope of this chapter.

- Place the neck in an extended position, using the shoulder roll.
- Give fentanyl for relaxation, prior to giving vecuronium.

Because of the risk of air embolism during the removal of the venous catheter, the infant must not be allowed to breathe during decannulation. If two doses of vecuronium do not produce paralysis, give pancuronium.

- Increase ventilator setting to a rate of 40 to 50 breaths/min, a peak inspiratory pressure of 20 to 25 cm H_2O (depending on chest movement), and F_iO_2 of 0.30 to 0.40 after paralytic agent is given.
- Clean the neck, and drape as for cannulation.
- Anesthetize with 0.25% Xylocaine with epinephrine.
- Cut and remove the Vicryl suture.
- Remove absorbable gelatin sponge packing, exposing the catheters and vessels.

If a jugular bulb catheter is in place, it is usually removed first to allow better visualization for removal of the venovenous ECMO catheter.

• The jugular bulb catheter should be clamped off before its removal, after the patient is taken off bypass. Be aware that removing the catheter while on bypass without a clamp in place will result in the introduction of air into the circuit.

In case of venoarterial ECMO, the venous catheter is usually removed first because it is most readily accessible.

- Separate the catheter from surrounding tissue by blunt dissection.
- Encircle the vein with a 2-0 silk tie, which is used for traction and hemostatic control.
- Place a Satinsky clamp around the vein to stabilize the catheter (Fig. 32.10).
- Place a 2-0 silk tie proximal to the clamp.
- Cut the silk ties securing the catheter in the vein with a no. 11 scalpel blade. The two proximal ties should be cut where they cross the vessel loop ("bootieâ€).
- Ask the ECMO specialist to remove the patient from the ECMO circuit.
- Monitor vital signs and oxygen saturation as an indication that ventilator settings are appropriate. Settings may have to be increased when the patient is removed from the circuit.
- Provide an inspiratory "hold†on the ventilator while the surgeon places pressure on the liver and removes the catheter. Failure to do this can result in air embolus.
- Replace any significant blood loss.
- Cut the 2-0 silk traction suture and tie the suture proximal to the Satinsky clamp. Remove the Satinsky clamp.
- Isolate the arterial catheter, dissect free, and remove.

The decannulation procedure is the same as for the arterial catheter, with the exception that an inspiratory hold is not required.

• Give protamine (1 mg IV) after removal of both catheters.

Administration of protamine is not mandatory if there is no significant bleeding.

- Irrigate the wound with sterile saline and cauterize any bleeding sites.
- If desired, pack the wound with a thrombin-soaked absorbable gelatin sponge and close the neck incision using subcuticular horizontal sutures of 4-0 Vicryl.
- Remove the sutures holding the cannula behind the right ear.
- Place povidone–iodine ointment over the incision and cover with semipermeable transparent dressing.



FIG. 32.10. Placement of Satinsky vessel clamp prior to removal of extracorporeal membrane oxygenation catheter.

F. Complications

- Vessel laceration, which may require a sternotomy for correction
- Excessive blood loss
- Venous air embolus

References

1. O'Rourke PP, Crone RK, Vacanti JP, et al. Extracorporeal membrane oxygenation and conventional medical therapy in neonates with persistent pulmonary hypertension of the newborn: a prospective randomized study. Pediatrics. 1989;84:957.

2. UK Collaborative ECMO Trial Group. UK Collaborative Randomised Trial of Neonatal Extracorporeal Membrane Oxygenation. Lancet. 1996;348:75.

3. Elbourne D, Field D, Mugford M. Extracorporeal membrane oxygenation for severe respiratory failure in newborn infants. Cochrane Database Syst Rev. 2002:CD001340.

4. Kanto WP, Shapiro MB. The development of prolonged extracorporeal circulation. In: Zwischenberger JB, Steinhorn RH, Bartlet RH, eds. Extracorporeal Cardiopulmonary Support in Critical Care. 2nd ed. Ann Arbor, MI: Extracorporeal Life Support Organization; 2000:27.

5. Rais-Bahrami K, Short BL. The current status of neonatal extracorporeal membrane oxygenation. Semin Perinatol. 2000; 24:406.

6. Bartlett RH, Roloff DW, Custer JR, et al. Extracorporeal life support: the University of Michigan experience. JAMA. 2000; 283:904.

7. Rais-Bahrami K, Van Meurs KP. ECMO for neonatal respiratory failure. Semin Perinatol. 2005;29:15. 8. Allison PL, Kurusz M, Graves DF, et al. Devices and monitoring during neonatal ECMO: survey results. Perfusion. 1990;5:193. 9. Van Meurs KP, Mikesell GT, Seale WR, et al. Maximum blood flow rates for arterial cannulae used in neonatal ECMO. ASAIO Trans. 1990;36:M679.

 Irish MS, O'Toole SJ, Kapur P, et al. Cervical ECMO cannula placement in infants and children: recommendations for assessment of adequate positioning and function. J Pediatr Surg. 1998;33:929.
 Neonatal ECMO Registry of the Extracorporeal Life Support Organization (ELSO). Ann Arbor, MI: ELSO; July 2005.

12. Rais-Bahrami K, Rivera O, Mikesell GT, et al. Improved oxygenation with reduced recirculation during venovenous extracorporeal membrane oxygenation: evaluation of a test catheter. Crit Care Med. 1995;23:1722.

13. Rais-Bahrami K, Waltom DM, Sell JE, et al. Improved oxygenation with reduced recirculation during venovenous ECMO: comparison of two catheters. Perfusion. 2002;17:415.

14. Paul JJ, Desai H, Baumgart S, et al. Aortic dissection in a neonate associated with arterial cannulation for extracorporeal life support. ASAIO. 1997;43:92.

33-Management of Vascular Spasm and Thrombosis

Jayashree Ramasethu

Thromboembolism (TE) is being increasingly recognized as a significant complication of intravascular catheters in sick newborn infants. About 90% of neonatal venous thromboses are associated with central venous catheters, although additional risk factors may be present (1). This chapter focuses on catheter-related vascular spasm and thrombosis.

A. Definitions

- Vascular spasm is transient, reversible arterial constriction, triggered by intravascular catheterization or arterial blood sampling. The clinical effects of vascular spasm usually last less than 4 hours from onset, but the condition may be difficult to differentiate from more serious thromboembolic (TE) disease. The diagnosis of vasospasm of arteries may only be made retrospectively, after documentation of the transient nature of ischemic changes and complete recovery of circulation.
- Thrombosis is the complete or partial obstruction of arteries or veins by blood clot(s).

B. Assessment

- Clinical diagnosis (Fig. 33.1)
 - The clinical signs associated with arterial or venous TE are shown in Table 33.1.
 - Vascular spasm of peripheral arteries is characterized by transient pallor or cyanosis of the involved extremity with diminished pulses and perfusion.
 - Persistent bacteremia or thrombocytopenia are nonspecific signs associated with vascular thrombosis at any site.
 - Clinical signs may be subtle or absent in many cases of thrombosis.
- Diagnostic imaging
 - Contrast angiography is the "gold standard.†It gives the best definition of thrombosis, but it is difficult to perform in critically ill neonates and requires infusion of radiocontrast material that may be hypertonic or cause undesired increase in vascular volume (2).
 - Doppler ultrasonography is portable, noninvasive, and can be used to monitor progress over time. It is useful for follow-up of aortic and peripheral arterial obstruction, but it may give both false positive and false negative results compared with contrast angiography (3).
- Additional diagnostic tests
 - Obtain detailed family history in all cases of unusual or extensive TE.

- In the absence of predisposing risk factor for thrombosis, consider investigations for the diagnosis of thrombophilic disorders: anticardiolipin, antithrombin III, protein C, protein S deficiency, etc. (4,5).
- C. Management of Vascular Spasm
 - Warm contralateral extremity (reflex vasodilation).
 - Maintain neutral thermal environment for affected extremity, i.e., keep heat lamps away from the area.
 - Maintain limb in horizontal position.
 - If necessary, restore circulating volume with blood, albumin, or plasma.
 - Consider removing catheter.

A

If mild cyanosis of the fingers or toes is noted after insertion of an arterial catheter, but peripheral pulses are still palpable, a trial of reflex vasodilation with close observation is reasonable, because vasospasm may resolve. Continually assess the need for keeping the catheter in place (i.e., the benefits of arterial access vs. the risk of thrombosis and further complications). A white or $\hat{a}\in \hat{c}$ appearing extremity is an indication for immediate removal of the catheter.

- Topical nitroglycerine has been demonstrated to reverse peripheral and umbilical artery catheter–induced ischemia in isolated case reports. Maintain good circulatory volume. Monitor for hypotension and be prepared to treat it immediately.
 - Topical application of 2% nitroglycerine ointment at a dose of 4 mm/kg body weight, applied as a thin film over the affected areas; may be repeated after 8 hours (6,7 and 8).
 - Application of a nitroglycerine patch has also been described (9).



FIG. 33.1. Skin necrosis associated with an umbilical artery catheter. Such lesions develop after vasospasm or embolization. A: Spinal injury may be present when ischemia involves this region. B: The distal part of an extremity is a common site for embolic arterial loss. The full extent of loss is unpredictable at this stage. (From Fletcher MA. Physical Diagnosis in Neonatolgy. Phildelphia: Lippincott-Raven; 1998:127)

	TABLE 33.1 Diagnosis of Vascular Thrombosis	
Site	Clinical Signs	Diagnostic Imaging
CVL-associated venous	Malfunction of CVL, SVC syndrome, chylothorax,	Contrast angiography
thrombosis	swelling and livid discoloration of extremity, dilatation of	Doppler
	collateral veins over trunk or abdomen in chronic cases	ultrasonography
Inferior vena cava	Lower limbs cool, cyanotic, edematous	Real time 2-D
thrombosis		ultrasonography
Superior vena cava	Swelling of upper limbs and head, chylothorax	
thrombosis		
Renal vein thrombosis	Flank mass, hematuria, thrombocytopenia, hypertension	
Aortic or renal arterial	Systemic hypertension, hematuria, oliguria	
thrombosis		
Peripheral or central (aorta	Pallor, coldness, weak or absent peripheral pulse(s),	
or iliac) arterial thrombosis	discoloration, gangrene	
Right atrial thrombosis	Congestive heart failure	Echocardiography
Pulmonary	Respiratory failure	Lung perfusion scan
thromboembolism		

CVL, central venous line; SVC, superior vena cava.

D. Management of Catheter-Related Thromboembolism

- General principles
 - Thrombolysis to restore catheter patency is described in Chapter 31.
 - Management of vascular TE may involve one or more of the following: supportive care, anticoagulation, fibrinolytic therapy, surgical intervention (4,5,10).
 - Treatment for neonates is highly individualized and is determined by the extent of thrombosis and the degree to which diminished perfusion to the affected extremity or organ affects function (4,5).
 - The infant should be managed in an appropriately staffed and equipped neonatal intensive care unit, where anticoagulant or thrombolytic therapy can be administered and monitored, and supportive and surgical care are readily available. Consultation with pediatric hematology is recommended. Plastic or vascular surgical consultation may be required.
 - The International Children's Thrombophilia Network, based in Canada, is a free consultative service, maintained 24 hours a day, for physicians worldwide who are caring for children with thromboembolic disease. The toll-free line in North America is 1-800-NO-CLOTS; the number for physicians elsewhere is 1-905-573-4795. The service provides current management protocols as well as links to the network and its services (11).
- Initial management
 - Initiate management as for vascular spasm for peripheral arterial ischemia.
 - Removal of catheter is advocated (5).

Peripheral arterial and umbilical catheters should be removed immediately if an extremity appears white or $\hat{a} \in \hat{c}$ blanched. $\hat{a} \in \hat{c}$ Intravascular catheters are left in place even in the presence of thrombosis only if local thrombolysis through the catheter is planned (12).

• Supportive care: correct volume depletion, electrolyte abnormalities, anemia, and thrombocytopenia; treat sepsis.

• Anticoagulant/thrombolytic therapy (4,5)

The risk of serious bleeding associated with antithrombotic therapy in neonates must be balanced against the possibility of organ or limb loss or death without appropriate treatment. Adequate randomized controlled trials to guide therapy in neonates are not available (12,13). The following guidelines are based on published protocols used in small case series.

- Contraindications
 - Major surgery within the last 10 days
 - Major bleeding: intracranial, pulmonary, or gastrointestinal
 - Pre-existing cerebral ischemic lesions
 - Relative contraindications: thrombocytopenia (platelet count <100 x 10⁹/L), hypofibrinogenemia (fibrinogen <100 mg/dL), severe coagulation factor deficiency, hypertension. Anticoagulant/thrombolytic therapy may be given after correction of these abnormalities.
- \circ Precautions
 - No arterial punctures
 - No subcutaneous or intramuscular injections
 - No urinary catheterizations
 - Avoid aspirin or other antiplatelet drugs.
 - Monitor serial head ultrasound scans for intracranial hemorrhage.
- o Anticoagulants
 - Oral anticoagulants (Warfarin)

Warfarin is not recommended in neonates because its anticoagulant effects are altered significantly by diet, illness, and other medications.

- Standard or unfractionated heparin (14,15)
 - Anticoagulant, antithrombotic effect limited by low plasma levels of antithrombin in neonates
 - Dosage: Loading dose of standard heparin 75 U/kg IV over 10 minutes; maintenance dose 28 U/kg/hr for infants <1 year of age
 - Monitoring: Check activated partial thromboplastin time (APTT) 4 hours after loading dose and 4 hours after every change in the infusion rate.
 - Adjust heparin infusion to maintain APTT at 60 to 85 seconds, corresponding to an anti-FXa level of 0.3 to 0.7 U/mL.
 - If APTT <60 seconds, increase infusion rate by 10%, and repeat APTT after 4 hours.
 - If APTT >85 seconds, decrease infusion rate by 10%, and repeat APTT after 4 hours.
 - If APTT >96 seconds, hold heparin infusion for 30 to 60 minutes, and restart at lower infusion rate; repeat APTT after 4 hours.
 - When APTT values are therapeutic and infusion rate is stable, obtain complete blood count and APTT daily.
 - The optimal duration of anticoagulation with heparin is unknown, but therapy is usually continued for 5 to 14 days (4,5).
 - Monitor thrombus closely both during and following treatment.
 - Complications

•

- Bleeding
- Heparin-induced thrombocytopenia (16)
- Antidote: Protamine sulfate IV may be used for rapid reversal of bleeding caused by heparin.

Dosage: 1.0 mg/100 U heparin received if the time since the last heparin dose is <30 minutes. The dose of protamine decreases linearly with increasing time since the last heparin dose, dropping to 0.25 mg/100 U heparin if the time since the last heparin dose is >120 minutes (5).

TABLE 33.2 Guidelines for Adjusting LMWH Treatment Dosage (32)

Anti-Factor		
Xa Level	Dose Change	Repeat Anti-Factor Xa Level
<0.35 U/mL	↑ by 25%	4 h following dose adjustment
0.35-0.49	↑ by 10%	4 h following dose adjustment
U/mL		
0.50-1 U/mL	No change	Weekly, 4 h following a dose ^a
1.1-1.5 U/mL	↓ by 20%	Before next dose and 4 h following dose
		adjustment
1.6-2.0 U/mL	Hold next dose for 3 h, then T by	Before next dose and 4 h following dose
	30%	adjustment
>2.0 U/mL	Hold dose until anti-factor Xa	Q 12 h until anti-factor Xa level <0.5, then 4
	level 0.5 U/mL, then T by 40%	h following reinstitution of therapy
^a Check level ⁴	⁴ h after next dose if there is a char	nge in renal function, addition of antibiotics,
or if any bleed	ling occurs.	

Modified from Andrew M, deVeber G. *Pediatric Thromboembolism and Stroke Protocols*. Hamilton, Ont.: B.C. Decker; 1999, with permission.

• Low-molecular weight heparin (LMWH) (17,18)

- LMWHs have specific activity against factor Xa and less activity against thrombin, so therapy is monitored by anti-FXa assay and not by APTT.
- Different LMWH preparations (e.g., enoxaparin, dalteparin, reviparin) differ in their molecular weights and dosage regimens.
- Advantages: Subcutaneous administration.
- Dosage: starting dose 1.5 mg/kg subcutaneously every 12 hours. Therapeutic dose range may vary from 0.95 to 3.5 mg/kg/12h.
- Monitoring
 - Adjust dose to maintain anti-factor Xa level between 0.5 and 1 unit/mL (Table 33.2).
 - Although LMWHs have more predictable pharmacokinetics in adults, in neonates, prematurity, rapid growth, and liver and kidney dysfunction make dosage less predictable. Frequent adjustment of the dose is required to attain target anti-FXa levels.
 - Draw blood sample for testing from fresh venipuncture. There must be no contamination from standard heparin, e.g., from an arterial line.
 - Check levels 4 hours after subcutaneous administration of LMWH on day 1 and day 2 of treatment.
 - If therapeutic, weekly check of anti-factor Xa levels is adequate.
- To discontinue anticoagulation, simply discontinue LMWH therapy. If an invasive procedure such as lumbar puncture is required, skip two doses of LMWH, and measure anti-factor Xa level prior to the procedure.
- If an immediate antidote is required, protamine is partially effective. Consult a hematologist. 1 mg of protamine neutralizes 100 U (1 mg) of Enoxaparin given within the last 3 to 4 hours. Protamine should be administered IV, over a 10-minute period, as rapid infusions may produce hypotension.

- Thrombolytic agents
 - Thrombolytic agents should be considered in the presence of extensive or severe thrombosis when organ or limb viability is at risk.
 - Agents work by enhancing the conversion of plasminogen to plasmin, which then proteolytically cleaves fibrin within the clot to fibrin degradation products.
 - Supplementation with plasminogen in the form of fresh frozen plasma enhances the thrombolytic effect (19).
 - Thrombi that have been present for several days may be resistant to thrombolysis. Failure rates may be as high as 50% (20).
 - The administration of heparin, either concomitantly or following thrombolytic therapy, has not been adequately evaluated in neonates.
 - A wide variety of dosage protocols have been used (12).
 - Streptokinase
 - Associated with allergic reactions in adults and older children; replaced largely by urokinase. Limited information in neonates.
 - Dosage regimens in neonates range from 50 U/kg/hr (directly into the thrombus) to intravenous infusions of 2,000 U/kg/hr (5,21,22).
 - Urokinase
 - Nonantigenic, but like streptokinase, has a low affinity for plasminogen bound to fibrin (clot-bound fibrin).
 - Human-derived urokinase has been replaced by the recombinant form in the United States.
 - Dosage: Loading dose 4,400 U/kg over 20 minutes, followed by continuous IV infusion of 4,400 U/kg/hr for 6 to 12 hours (4,5). The dosage and duration of therapy is determined by the response. Doses up to 10,000 to 16,000 IU/kg/hr have been used (23). Smaller doses have been used for local treatment of thrombi without removal of central venous catheters.
 - Recombinant tissue plasminogen activator (r-tPA)
 - Nonantigenic, has specific affinity for plasminogen bound to fibrin, intensifying thrombolysis at the site of the clot; has a short half-life.
 - Large number of case reports published, with varying dosage protocols (17)
 - Dosage protocol 1: Infusion, 0.1 mg/kg/hr. If no response after 6 hours, may increase infusion rate by 0.1 mg/kg/hr every 6 hours, to a maximum of 0.5 mg/kg/hr. Monitor fibrinogen levels every 4 hours and keep levels >100 mg/dL (24).
 - Dosage protocol 2: Initial bolus, 0.1 mg/kg over 10 minutes, followed by infusion of 0.3 mg/kg/hr over a 3-hour period. Perform Doppler sonography at the end of each infusion. Fibrinogen assay 1 hour and 4 hours after each r-TPA infusion is initiated. If repermiabilization is not complete, up to four additional r-TPA infusions may be given at intervals of 12 to 24 hours (25).
 - Catheter-directed thrombolysis: Infusion of low doses of r-tPA through a catheter with the tip adjacent to or within the thrombus; higher response rate with decreased risk of bleeding (12).

Initial bolus dose ranges from 0 to 0.5 mg/kg, followed by infusion of 0.015 to 0.2 mg/kg/hr.
- Monitoring
 - Imaging studies every 4 to 12 hours during fibrinolytic therapy, to allow discontinuation of treatment as soon as clot lysis is achieved.
 - Measure thrombin time, fibrinogen and plasminogen levels, and fibrin split products or d-dimers prior to therapy, 3 to 4 hours after initiation of fibrinolytic therapy and one to three times daily thereafter (2).
 - Fibrinolytic response is measured by a decrease in fibrinogen concentration and increase in levels of fibrin degradation products, but the correlation between these hemostatic parameters and efficacy of thrombolysis is poor. Maintain fibrinogen levels of at least 100 mg/dL to prevent bleeding.
- E. Complications of Anticoagulation/Fibrinolytic Therapy
 - Hemorrhagic complications (12,26)
 - Intracerebral hemorrhage: Incidence approximately 1% in term neonates, 13% in preterm neonates, increasing to 25% in preterm infants treated in the first week of life. Data in preterm infants are confounded by the risk of "spontaneous†intraventricular hemorrhage (26).
 - Other major hemorrhages: Gastrointestinal, pulmonary
 - Bleeding from puncture sites and recent catheterization sites
 - Hematuria
 - Embolization

Dislodgement of intracardiac thrombus, causing obstruction of cardiac valves or main vessels, pulmonary or systemic embolization (27,28).

F. Surgical Intervention (10,29,30and 31)

Early consultation is recommended, because surgical management may be required concomitantly, particularly for life- or limb-threatening emergencies.

- Thrombectomy
- Microvascular reconstruction
- Leech therapy
- Decompressive fasciotomy
- Mechanical disruption of thrombus using soft wires and balloon angioplasty in conjunction with continuous site-directed thrombolytic infusion into the clot.
- Amputation

References

1. Chalmers EA. Epidemiology of venous thromboembolism in neonates and children. Thrombosis Res. 2006;118:3–12.

 Schmidt B, Andrew M. Report of the Scientific and Standardization Subcommittee on Neonatal Hemostasis. Diagnosis and treatment of neonatal Thrombosis. Thromb Hemostat. 1992;67: 381–382.
 Roy M, Turner-Gomes S, Gill G, et al. Accuracy of Doppler ultrasonography for the diagnosis of Thrombosis associated with umbilical venous catheters. J Pediatr. 2002;140:131–134.

4. Williams MD, Chalmers EA, Gibson BE; Haemostasis and Thrombosis Task Force, British Committee for Standards in Haematology. The investigation and management of neonatal haemostasis and Thrombosis. Br J Haematol. 2002;119: 295-309.

5. Monagle P, Chan A, Chalmers E, Michelson AD. Antithrombotic therapy in children. The Seventh ACCP Conference on Antithrombotic and Thrombolytic therapy. Chest. 2004;126: 645S–687S.

6. Wong AF, McCulloch LM, Sola A. Treatment of peripheral tissue ischemia with topical nitroglycerine ointment in neonates. J Pediatr. 1992;121:980–983.

7. Vasquez P, Burd A, Mehta R, et al. Resolution of peripheral artery catheter induced ischemic injury following prolonged treatment with topical nitroglycerine ointment in a newborn: a case report. J Perinatol. 2003;23:348–350.

8. Baserga MC, Puri A, Sola A. The use of topical nitroglycerine ointment to treat peripheral tissue ischemia secondary to arterial line complications in neonates. J Perinatol. 2002;22:416–419.

9. Varughese M, Koh THHG. Successful use of topical nitroglycerine in ischemia associated with umbilical arterial line in a neonate. J Perinatol. 2001;21:556–558.

10. Coombs CJ, Richardson PW, Dowling GJ, et al. Brachial artery Thrombosis in infants: an algorithm for limb salvage. Plast Reconstr Surg. 2006;117:1481–1488.

11. Andrew M. Society for Pediatric Research Presidential Address 1998: The SPR and 1-800-NO-CLOTS: a common vision. Pediatr Res. 1998;44:964–973.

12. Albisetti M. Thrombolytic therapy in children. Thromb Res. 2006;118:95–105.

13. John CM, Harkensee C. Thrombolytic agents for arterial and venous thromboses in neonates. Cochrane Database Syst Rev. 2005;25:CD004342.

14. Sutor Ah, Massicote P, Leaker M, Andrew M. Heparin therapy in pediatric patients. Semin Thromb Hemost. 1997;23:303–319.

15. Streif W, Mitchell LG, Andrew M. Antithrombotic therapy in children. Curr Opin Pediatr. 1999;11:56–64.

16. Kumar P, Hoppensteadt DA, Prechel MM, et al. Prevalence of heparin-dependent platelet activating antibodies in preterm newborns after exposure to unfractionated heparin. Clin Appl Thromb Hemost. 2004;10:335–339.

17. Ramasethu J. Management of vascular Thrombosis and spasm in the newborn. Neoreviews. 2005;6:e1–e14.

18. Streiff W, Goebel G, Chan AKC, Massicotte MP. Use of low molecular weight heparin (enoxaparin) in newborn infants: a prospective cohort study of 62 patients. Arch Dis Child Fetal Neonatal Ed. 2003;88:F365–F370.

19. Andrew M, Brooker L, Paes B, et al. Fibrin clot lysis by thrombolytic agents is impaired in newborns due to a low plasminogen concentration. Thromb Hemost. 1992;68:325–330.

20. Wever MLG, Liem KD, Geven WB, Tanke RB. Urokinase therapy in neonates with catheter related central venous Thrombosis. Thromb Haemost. 1995;73:180–185.

21. Cheah FC, Boo NY, Rohana J, Yong SC. Successful clot lysis using low dose of streptokinase in 22 neonates with aortic thromboses. J Paediatr Child Health. 2001;37:479–482.

22. Richardson R, Applebaum H, Touran T, et al. Effective thrombolytic therapy of aortic Thrombosis in the small premature infant. J Pediatr Surg. 1988;23:1198–1200.

23. Hausler M, Hubner D, Hornchen H, et al. Successful thrombolysis of inferior vena cava Thrombosis in a preterm neonate. Clin Pediatr. 2000;40:105–108.

24. Weiner GM, Castle VP, DiPietro MA, Faix RG. Successful treatment of neonatal arterial thromboses with recombinant tissue plasminogen activator. J Pediatr. 1998;133:133–136.

25. Farnoux C, Camard O, Pinquier D, et al. Recombinant tissue-type plasminogen activator therapy of Thrombosis in 16 neonates. J Pediatr. 1998;133:137–140.

26. Zenz W, Arlt F, Sodia S, Berghold A. Intracerebral hemorrhage during fibrinolytic therapy in children. A review of literature of the last thirty years. Semin Thromb Hemost. 1997;23: 321–332.

27. Nowak-Gottl U, Auberger K, Halimeh S, et al. Thrombolysis in newborns and infants. Thromb Haemost. 1999;82(suppl): 112–116.

28. Alkalay AL, Mazkereth R, Santulli T, et al. Central venous line Thrombosis in premature infants: a case management and literature review. Am J Perinatol. 1993;10:323–326.

29. Friedman J, Fabre J, Netscher D, Jaksic T. Treatment of acute neonatal vascular injuriesâ€"the utility of multiple interventions. J Pediatr Surg. 1999;34:940â€"945.

30. Robinson A, Fellows KE, Bridges ND, Rome JJ. Effectiveness of pharmacomechanical thrombolysis in infants and children. Am J Cardiol. 2001;87:496–499.

31. Blank JE, Dormans JP, Davidson RS. Perinatal limb ischemia: orthopaedic implications. J Pediatr Orthop. 1996;16:90–96.

32. Andrew M, deVeber G. Pediatric Thromboembolism and Stroke Protocols. Hamilton, Ont: BC Decker; 1999.

34-Endotracheal Intubation

Khodayar Rais-Bahrami

A. Indications

- When prolonged positive-pressure ventilation is required
- To relieve critical upper airway obstruction (Fig. 34.1)
- To provide a route for selective bronchial ventilation
- When tracheal suctioning is required to obtain direct tracheal culture
- To assist in bronchial hygiene when secretions cannot be cleared
- When diaphragmatic hernia is suspected

B. Contraindications

There is no absolute contraindication to intubating a neonate who has one of the above-mentioned indications. In older patients, the presence of cervical injuries is a contraindication to intubation with a laryngoscope; however, because the occurrence of cervical injuries is infrequent in neonates, we consider that endotracheal intubation is associated with less risk than performance of an emergency tracheotomy. C. Equipment

The supplies and equipment necessary to perform endotracheal intubation should be kept together on either a resuscitation cart or an intubation tray. Each delivery room, nursery, and emergency room should have a complete set of the following items.

- Humidified oxygen/air source, blender, and analyzer
- Oxygen tubing
- Resuscitation bag and mask
- Suctioning device, with 10-French (Fr) suction catheters
- Cardiorespiratory monitor
- Pulse oximetry oxygen saturation monitor
- Stethoscope
- Gloves
- Scissors
- Sterile stylet
- Adhesive tape: Two 8- to 10-cm lengths of $\hat{A}^{1/2}$ -in-wide tape, with half the length split and one 10- to 15-cm length unsplit
- Magill forceps (optional for nasotracheal intubation)
- Pediatric laryngoscope with an extra set of batteries and extra bulb
 - o Miller blade size 1 for full-term infant
 - Miller blade size 0 for preterm infant

Straight rather than curved blades are preferred for optimal visualization.

- Modified blade to allow continuous flow of oxygen at 1 to 2 L/min for better maintenance of oxygenation during procedure (1)
- Endotracheal tubes with internal diameters of 2.5, 3.0, 3.5, and 4.0 mm
 - Diameter selected for infant size (Table 34.1)
 - Length selected for infant's size (2,3 and 4)

In neonates, there is little leeway between a tube that is too high (increased risk for extubation) or too low (increased risk for mainstem intubation or airway trauma). The appropriate length for an endotracheal tube depends on a number of factors, including an infant's size, and it can be quickly and accurately estimated by measuring the nasal–tragus length (NTL) and/or sternal length (STL). The modified prediction formula for insertion by the orotracheal route is NTL or STL + 1. For the nasotracheal route, the formula is NTL or STL + 2 (5).

It is rarely necessary to insert a tube more than 1 to 2 cm below the vocal cords, regardless of the infant's size. Exceptions include the presence of anatomic defects that necessitate a $\hat{a}\in \hat{c}$ by pass $\hat{a}\in \hat{c}$ airway, such as a tracheal fistula or subglottic obstruction, and when selective bronchial intubation is intended (6). See Appendix D.

- D. Precautions (Table 34.2)
 - Select orotracheal route for all emergency intubations or when a bleeding diathesis is present. Reserve nasotracheal intubation for elective procedures after stabilization with orotracheal tube, unless oral anatomy precludes oral intubation.
 - Prepare all equipment before starting procedure. Keep equipment ready at bedside of patients likely to require intubation.
 - Use appropriate-size tubes (Table 34.1). The tube should not fit tightly between the vocal cords to minimize upper airway trauma.
 - To minimize hypoxia, each intubation attempt should be limited to 20 seconds. Interrupt an unsuccessful attempt to stabilize the infant with bag-and-mask ventilation. In most cases an infant can be adequately ventilated by bag and mask, so endotracheal intubation can be achieved as a controlled procedure. The one important exception is in a case of prenatally diagnosed or suspected congenital diaphragmatic hernia.
 - Recognize anatomic features of neonatal upper airway (Fig. 34.2).
 - Ensure visualization of larynx. This is the most important step (Fig. 34.3).
 - Have assistant maintain proper position of patient.
 - o Avoid hyperextending or rotating neck.





FIG. 34.1. A: Vallecula cyst, causing stridor and proximal airway obstruction. B: Endotracheal tube passes beneath cyst. C: Same patient after laser surgical treatment.

TABLE 34.1 Endotracheal Tube Diameter for Patient Weight and Gestational AgeTube Size (ID mm)WeightGestational Age

2.5	<1,000 g	<28 wk
3.0	1,000–2,0	00 g28–34 wk
3.5	2,000–3,0	00 g34–38 wk
4.0	>3,000 g	>38 wk

Problem	Suggested Approach for Solution	
Infant's tongue gets in way.	Push tongue aside with finger before inserting blade.	
Secretions prevent visualization.	Suction prior to intubation attempt.	
Tube seems too big to fit through	Verify correct tube size for patient weight and gestational age.	
vocal cord.		
Vocal cords are closed.	Decrease angle of neck extension.	
	Apply traction to blade.	
	Select smaller tube size.	
	Evaluate for airway stenosis.	
Unsure of appropriate tube	Await spontaneous breath.	
length.	Apply gentle suprasternal pressure.	
Difficult to ventilate after	Insert tube just past vocal cord.	
intubation.	Predetermine tube length.	
	Obtain chest radiograph with head in neutral position to confirm	
	tube position relative to carina.	
Swelling of neck and anterior	Verify that tube is in trachea.	
chest.	Verify that tube is not in bronchus.	
	Consider tube and/or airway obstruction.	
Blood return from endotracheal tube.	Evaluate for tracheal perforation.	
Tube slips into main bronchus.	Avoid neck extension.	
	Secure tape fixation.	
	Maintain correct lip-to-tip distance.	
Unplanned extubation.	Regularly verify correct tube distance.	
	Secure tape and replace as necessary.	
	Support neck when moving infant.	
	Avoid neck extension or traction on tube.	
	Secure infant's hands.	

TABLE 34.2 Trouble-Shooting Problems with Endotracheal IntubationSuggested Approach for Solution



FIG. 34.2. Anatomic view of neonatal upper airway. The glottis sits very close to the base of the tongue, so visualization is easiest without extending the neck.

- Do not use pressure or force that may predispose to trauma.
 - Avoid using maxilla as fulcrum for laryngoscope blade.
 - Avoid excessive external tracheal pressure.
 - Avoid pushing tube against any obstruction.



FIG. 34.3. A: Normal epiglottis obscuring glottis. This amount of clear secretions does not require suctioning for visualization. B: Same airway as in Fig. 34.1 after surgical removal of cyst. Glottic opening is visible just beneath epiglottis. Gentle tracheal pressure or decreasing neck extension while lifting tip of laryngoscope blade will improve visibility.

- Make certain all attachments are secure.
 - Avoid obscuring the point of connection of tube and adapter with any fixation device.
 - Secure tube carefully in position to avoid dislodgement, kinking, or movement.
 - Vary contact point from side to side to prevent damage to developing palate and palatal ridges (7,8,9 and 10).
 - Note relationship of head position to intratracheal depth of tube on radiograph (11,12).
- Do not leave endotracheal tube unattached from continuous positive airway pressure; the natural expiratory resistance is lost by bypassing the upper airway.
- Recognize that in neonates, endotracheal tubes are often pushed in too far because of the short distance from the glottis to the carina. Use a standardized graph or location device (2,5).
- Recognize the association of short trachea (fewer than 15 tracheal cartilage rings) with certain syndromes: DiGeorge syndrome, skeletal dysplasias, brevicollis, congenital rubella syndrome, interrupted aortic arch, and other congenital syndromes involving the tracheal area (13).
- Identify and prevent the factors that are most likely to contribute to spontaneous extubation (14).
 - Increased secretions
 - Necessitating more frequent suctioning
 - Loosening of tape
 - Infant activity
 - Procedures requiring repositioning infant
 - Tube slippage

E. Technique (See also Endotracheal Intubation on the Procedures DVD, and Appendix D for Techniques of

Intubation Specific to Unique Patient Needs)

Orotracheal Intubation (Table 34.2)

• Position infant with the head in midline and the neck slightly extended, pulling chin into a "sniff†position (Fig. 34.4). The head of the infant should be at operator's eye level.

It may be helpful to place a roll under the baby's shoulders to maintain slight extension of the neck.

- Put on gloves.
- Clear oropharynx with gentle suctioning.
- Empty stomach.
- Bag-and-mask ventilate and preoxygenate infant as indicated by clinical condition. Follow heart rate and oxygenation.



FIG. 34.4. Appropriate sniff position for intubation. Note that the neck is not hyperextended; the roll provides stabilizing support.

- Turn on the laryngoscope light, and hold the laryngoscope in left hand with thumb and first three fingers, with the blade directed toward patient.
 - Put thumb over flat end of laryngoscope handle.
 - Stabilize the infant's head with right hand.

The laryngoscope is designed to be held in the left hand, by both right- and left-handed individuals. If held in the right hand, the closed, curved part of the blade may block the view of the glottis as well as make insertion of the endotracheal tube impossible.



FIG. 34.5. Open the mouth and push the tongue aside with the forefinger while stabilizing the head with the thumb and other fingers of the right hand.

- Open infant's mouth and depress tongue toward the left with the back of right forefinger (Fig. 34.5).
 - Continue to steady head with rest of right hand.
 - Do not use the laryngoscope blade to open mouth.
- Under direct visualization, insert laryngoscope blade, sliding over the tongue until the tip of the blade is resting in the vallecula (the area between the base of the tongue and the epiglottis) (Fig. 34.6).

In general, the blade tip should be placed in the vallecula. However, in extremely premature infants, the vallecula may be too small, in which case it may be necessary to use the blade tip to gently lift the epiglottis.



FIG. 34.6. Pass the laryngoscope carefully along the finger to the back of the oropharynx.

• Lift the laryngoscope blade to open mouth further and simultaneously tilt the blade tip slightly to elevate the epiglottis and visualize the glottis (Fig. 34.7).

When lifting the blade, raise the entire blade in the direction that the handle is pointing. Do not lift the tip of the blade by using the upper gum line as the fulcrum for a rocking motion; this will not produce a clear view of the glottis and will place excessive pressure on the alveolar ridge, potentially impeding future tooth formation.



FIG. 34.7. With the laryngoscope at the proper depth, tilt the blade with the tongue as the fulcrum; at the same time, pull on the laryngoscope handle to move the tongue without extending the infant's neck. Use more traction than leverage.

- Suction as necessary.
- Have an assistant apply gentle pressure at the suprasternal notch to open the larynx and to feel the tube pass (15).
- Hold tube in right hand with concave curve anterior, and pass it down the right side of the mouth, outside the blade, while maintaining direct visualization (Fig. 34.8).



FIG. 34.8. Visualize the glottis and pass the endotracheal tube into the oropharynx. Keep the tube outside the curve of the laryngoscope blade for better mobility.

- Once the vocal cords and trachea are visualized, insert the endotracheal tube through vocal cords, approximately 2 cm into trachea or until the tip is felt passing the suprasternal notch by the assistant (Fig. 34.9).
- If the tube appears too large or does not pass easily, decrease angle of neck extension.
- Confirm endotracheal tube position within the trachea (16,17).
 - We currently use Pedi-Cap (Nellcore, Waukesha, WI) end-tidal CO_2 detector to verify the position of the endotracheal tube within the trachea. This technique responds quickly to exhaled CO_2 with a simple color change from purple to yellow. It also features an easy-to-see display window that provides constant visual feedback with breath-to-breath response (Fig. 34.10).
 - While gently ventilating with an Ambu bag, auscultate to make sure the breath sounds and chest movement are equal in both sides of the chest.
 - Observe respiratory wave pattern on oscilloscope to determine that artificial breath is at least as effective as spontaneous breath.
 - Verify lip-to-tip distance.



FIG. 34.9. A: Pass the endotracheal tube through the glottis to the appropriately predetermined length and remove laryngoscope. B: An assistant applies gentle pressure in the suprasternal notch to open the larynx and to detect when the tube passes into the trachea.

- If the endotracheal tube is correctly placed in the midtracheal region, there should be
 - Pedi-Cap response to exhaled CO₂ by a reversible color change, purple to yellow
 - Equal bilateral breath sounds
 - Slight rise of the chest with each ventilation



FIG. 34.10. Pedi-Cap CO₂ detector. Pedi-Cap is a trademark of Tyco Healthcare Group LP. (Reprinted by permission from Nellcor Puritan Bennett, Inc).

- No air heard entering stomach
- No gastric distention
- Suction endotracheal tube with sterile catheter, following technique described under F.
- Attach appropriate mechanical ventilatory device.
- Adjust required F_iO₂.
- Secure the tube to the infant's face (Figs. 34.11 and 34.12).

When using adhesive tape, make sure that the face is dried thoroughly to ensure adherence of the tape and to protect the skin. A more permanent fastening can be done later when a radiograph confirms correct placement of the endotracheal tube (18).

• Obtain chest radiograph with head in neutral position, and note the lip-to-tip distance and direction of bevel (Figs. 34.13 and 34.14).

When a correct tube length has been determined for the infant, note the tube marking at the level of the infant's lips.

• Cut off excess tube length to leave 4 cm from the infant's lips and reattach adapter firmly.

If a longer external length is required, before replacing the adapter, slip a short length of a larger endotracheal tube around the narrower tube to prevent kinking, for example, a 6-cm length of 3.5-mm tube over a 2.5-mm tube.

- Reconfirm tube marking at lip regularly, to avoid unnoticed advancement of the tube into the airway.
- Retape tube as necessary to maintain stability.

Nasotracheal Intubation

In neonates, orotracheal intubation is preferred because it is easier and faster to perform and there are few proven advantages to nasal intubations in small infants (19,20). Nasotracheal tubes are preferred in very active infants with copious oral secretions, making it difficult to keep the tube taped in position. When

anatomy precludes oral intubation or for oral surgery, nasotracheal intubation may become necessary. Premedication with succinylcholine/atropine or morphine/atropine has been reported to allow a shorter time for intubation, with fewer negative systemic effects (21).



FIG. 34.11. After initially determining that the endotracheal tube is in the correct position, connect the tube to an artificial ventilation source. In the term neonate begin fixation of the tube by painting the philtrum with tincture of benzoin and allowing it to dry. Avoid use of tincture of benzoin in low-birthweight infants; it increases epidermal stripping.



FIG. 34.12. Fixation of tube with half split tape. A: First half of one split tape (1) encircles the tube, and the other half (2) attaches to the upper lip. B: Second split half (3) attaches to the upper lip, while the bottom half (4) encircles the tube.

• Use sterile endotracheal tube. If stylet is used to curve tube, remove it prior to nasal insertion.



FIG. 34.13. Although the carina is usually at the level of T4 on the anteroposterior supine chest radiograph, this relationship may be significantly disturbed by a number of factors, including radiographic technique (x-ray tube position, angulation). For this reason and because the carina is usually easily visualized, as in these cases, one should directly relate the tip of the endotracheal tube to the carina radiographically, knowing the position of the head at the time of film exposure. In both cases, films were taken to verify endotracheal tube position but demonstrated problems with other procedures. A: Appropriate radiographic angle. (Note the oral gastric tube in the esophagus and not reaching the stomach.) B: Slightly lordotic radiographic angle. (Note the central venous line coiled in the heart.)



FIG. 34.14. Sequential radiographs demonstrate the effect of head rotation on bevel direction. A: With the head rotated to the right, the bevel appears to be directed against the tracheal wall. B: The head is rotated to the left, and the bevel is now positioned properly. If the bevel is directed against the posterior tracheal wall in a spontaneously breathing infant, there may be symptoms of tracheal obstruction on expiration. Rather than turning the head to achieve satisfactory position, rotate the endotracheal tube and retape in position.

- If desired, premedicate with atropine (20 mcg/kg) and succinylcholine (2 mg/kg) just before inserting tube. Be prepared to provide assisted bag-and-mask ventilation.
- If orotracheal tube is already in place, release fixation and position at far left of the mouth, to allow continued ventilation during nasotracheal intubation.
- Directly visualize oropharynx with laryngoscope as described previously, taking particular care not to hyper-extend neck.
- Suction oropharynx while keeping laryngoscope in place.
- Insert tube through nostril following natural curve of nasopharynx.

- As tube passes into the pharynx, align the tip with the center of the tracheal orifice, moving infant's head as needed.
- When the tip of the nasotracheal tube appears to be in direct line with the glottis, have assistant carefully withdraw the orotracheal tube.
- Apply gentle pressure over the suprasternal notch and advance tube through cords.

Use of the Magill forceps is often more cumbersome than helpful in smaller infants. A Magill forceps should always be available, but in a properly positioned infant, a curved tube usually passes directly into the trachea without forceps unless the neck is excessively extended, flexed, or rotated. Secure tube and verify position. The length of a nasotracheal tube for correct positioning of the tip in the trachea is approximately 2 cm longer than the equivalent length of an orotracheal tube.

F. Tracheal Suctioning

Suctioning of the nose, mouth, and pharynx is potentially quite traumatic in neonates. The same equipment, precaution, and complications apply as for tracheal suctioning. Always suction an endotracheal tube before suctioning the mouth; suction the mouth before the nose.

- Indications
 - To clear tracheobronchial airway of secretions
 - To keep artificial airway patent
 - To obtain material for analysis or culture
- Relative contraindications
 - Recent surgery in the area
 - Extreme reactive bradycardia
 - Pulmonary hemorrhage
 - Oscillatory ventilation
- Equipment

Sterile

- Saline for instillation into airway
- Saline or water for irrigation of catheter
- Gloves
- Suction catheters
 - Available safety features
 - Markings at measured intervals
 - Microscopically smooth surface
 - Multiple side holes in different planes
 - Large-bore hole for occlusion to initiate vacuum
 - No more than half the inside diameter of artificial airway
 - Use 8 Fr for endotracheal tube >3.5 mm.
 - Use 5 Fr for endotracheal tube <3.5 mm.
- Modified endotracheal tube adapter that allows passage of suction catheter without disconnecting tube from ventilator (Novometrix C/S Suction Adapter; Novometrix Medical Systems, Wallingford, CT, USA) (22)

- Vacuum source
 - Pressure set just high enough to move secretions into suction catheter
 - Mechanically controlled pressure source Pressure generated by oral suction on mucus extractors can be extremely variable and dangerously high (23).

Nonsterile

- Adjustable vacuum source with specimen trap, tubing, and pressure gauge
- Ventilatory device as indicated
 - o Manometer
 - Warmed, humidified oxygen at controlled level
 - Bag with positive end-expiratory pressure device
- Precautions
 - When feasible, use two people when suctioning airway, to minimize risk of patient compromise and complications and to shorten time for procedure.
 - Determine for each patient if it is better to continue mechanical ventilation during suctioning or to use a sigh with inflation hold after suctioning. Consider effect of interruption of ventilator therapy and loss of lung volume with each catheter passage.
 - Allow patient to recover between passages of catheter.
 - \circ $\;$ Stabilize head and airway to prevent tube dislodgement.
 - Assess secretions by auscultation and palpation to determine frequency for suctioning.
 - Avoid unnecessary suctioning just to follow a schedule.
 - Schedule prophylactic suctioning for tube patency only as often as needed to maintain it.
 - Consider increase in monitored airway resistance as indication for suctioning (24).
 - Readjust humidification as indicated by catheter and volume of secretions.
 - Avoid inadvertent suction during insertion of catheter.

Use lowest vacuum pressure effective in clearing secretions within a few seconds.

- Do not insert catheter as far as it will go or until reflex cough occurs. Use prescribed length.
 Do not suction if catheter is inserted too far; just touching the catheter to the tracheal wall may cause trauma.
- Limit time of insertion and suctioning to least time required to remove secretions.
- Technique for intubated patients
 - For artificial airways, use sterile technique with one sterile gloved hand and one free hand.
 - Monitor oxygen saturation continuously during suctioning.
 - Monitor heart rate continuously.
 - It is usually best to remove infant from ventilator and have second person perform assisted ventilation manually, using following guidelines adjusted to individual needs:
 - F_iO_2 set at or up to 10% higher than baseline
 - Monitor oxygenation. Adjust F_iO₂ to prevent swings in oxygenation.
 - Evaluate effect of procedure.
 - Peak inspiratory pressure as on ventilator or up to 10 cm H₂O higher
 - Continuous distending airway pressures same as on ventilator
 - Respiratory rate 40 to 60 breaths/min, applying an inspiratory hold intermittently

When there is a high risk of pulmonary air leak as in the presence of significant interstitial emphysema, it may be safer to use a technique of rapid manual ventilation at lower peak pressure instead of sighing with a prolonged inspiratory pressure. In other cases in which loss of lung volume with suctioning is of greater concern, use

sigh with a hold on inflation at a rate similar to ventilator. With suctioning, there is a loss of lung volume with a decrease in compliance. The adverse effect persists for a significant time when mechanical ventilation at the same setting is used during and after the suction procedure.

- Determine length of endotracheal tube plus adapter and note on suction catheter as limit of depth of insertion.
- Set vacuum at lowest level to achieve removal of secretions. The level of vacuum required depends on a number of variables, including
 - Air tightness of system and fluctuations in generated vacuum pressure
 - Accuracy of manometer
 - Diameter of catheter (smaller catheter, higher pressure)
 - Thickness and tenacity of secretions
- Holding catheter in one hand, moisten tip with water or saline. Note appropriateness of suction level by rate of liquid uptake. Adjust pressure with free hand.
- Open artificial airway with free hand.
 - Detach from bag; hold oxygen near end of tube, or
 - Open suction port of specialized endotracheal tube adapter.
- With free hand, stabilize airway. Pass catheter down airway to depth limit noted for the patient's endotracheal tube. Do not apply vacuum during insertion (i.e., keep suction control port open).
- Close proximal suction control port and withdraw catheter.
- Limit time for insertion and removal to 15 to 20 seconds.
- Reattach endotracheal tube to bag and ventilate for 10 to 15 breaths or until patient is stable.
 - Note oxygenation.
 - Note heart rate.
 - Note chest excursions.
- If secretions are thick or tenacious, instill 0.25 mL of saline into endotracheal tube and continue ventilation.
- Clear catheter with sterile water.
- Repeat process until airway is clear.

G. Fixation Techniques

Many fixation devices and techniques have been described in the literature. None of them can prevent all accidental extubations or malpositions (18,25,26 and 27). Here, we describe a simple and effective method.

- Prepare two 8- to 10-cm lengths of adhesive tape split half of the length and one 10- to 15-cm length without a split.
- Paint skin adjacent to the sides of the mouth and above the lips with tincture of benzoin (Fig. 34.11).
- Allow to dry while holding the tube in place.
- Tape the unsplit end of the adhesive to the cheek on one side of the mouth, and wrap the bottom half clockwise around the endotracheal tube just at the lip. Fold the last 2-mm end of tape on itself to leave a tab for easier removal (Fig. 34.12).
- Secure the other half above the upper lip.
- Repeat the procedure from the other side, reversing the direction of the taping and securing half on that side of the upper lip (Fig. 34.12).
- Secure one end of the long tape to one cheek at the zygoma. Loop the tape around the tube, and secure the other end to a similar point on the opposite cheek.
- Note the markings on the endotracheal tube at the level of the lips and the tape.
- Whenever the tape seems to be loosened from secretions, remove tape and repeat application of benzoin while holding tube at proper lip-to-tip depth.

H. Planned Extubation

Various vasoconstrictors and anti-inflammatory medications have been recommended to reduce postextubation stridor and to improve the success of extubation. Systemically administered dexamethasone appears to have very little if any effect in reducing acute postextubation stridor in neonates and children (28,29). Local application of steroids directly to the vocal cords has not been well studied.

- Perform chest physiotherapy and suction prior to extubation.
- Release all fixation devices while holding tube in place.
- Using manual ventilation, give infant a sigh breath, and withdraw tube during exhalation.
- Avoid suctioning during tube withdrawal, unless specifically removing foreign material from trachea.
- Allow recovery time before suctioning oropharynx.
- Keep the inspired gases well humidified.



FIG. 34.15. Subglottic erosion and stenosis after intubation.

I. Complications

- Acute trauma (30,31,32 and 33)
 - Tracheal or hypopharyngeal perforation
 - Pseudo-diverticulum
 - Hemorrhage
 - Laryngeal edema
 - Mucosal necrosis (Fig. 34.15)
 - Vocal cord injuries
 - Dislocation of arytenoid
- Chronic trauma (31,32,33,34,35,36,37, and 38)
 - Cricoid ulceration and fibrosis
 - Glottic and/or subglottic stenosis (Fig. 34.15)
 - Subglottic granuloma (Figs. 34.16 and 34.17)
 - Hoarseness, stridor, wheezing
 - Subglottic cyst
 - o Tracheomegaly
 - Protrusion of laryngeal ventricle
- Interference by oral tube with oral development (7,8,9 and 10,39,40,41,42, and 43)
 - Alveolar grooving
 - Palatal grooves (Fig. 34.18)
 - Acquired oral commissure defect (Fig 34.19)



FIG. 34.16. Radiographic magnification high-kilovoltage film (2 x) demonstrating an abrupt cutoff of the right bronchus intermedius (arrow) due to an endobronchial granuloma with secondary volume loss at the right lung base. Although these granulomas may be due to endotracheal tube trauma, in this area they are more likely related to suction tube injury. The endotracheal tube is just entering the right bronchus.



FIG. 34.17. Glottic granuloma after intubation. Epiglottis is manually retracted to reveal granuloma below cords. Esophageal opening is clearly visible beneath airway.



FIG. 34.18. Palatal groove after prolonged oral intubation. Such grooves may be seen after prolonged use of endotracheal or oral gastric tubes when the normal forces of the tongue are prevented from assisting palatal development.



FIG. 34.19. Acquired oral commissure defect: a complication of prolonged endotracheal intubation. (Reprinted by permission from Macmillan Publishers Ltd. Journal of Perinatology. 2005;25:612.)



FIG. 34.20. Nasal stenosis due to nasal cartilage necrosis following an indwelling nasotracheal tube.

- Posterior cross-bite
- Defective dentition

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- Enamel hypoplasia
- Incisor hypoplasia
- Poor speech intelligibility
- Local effects from nasal tube (44,45 and 46)
 - Erosion of nasal septum
 - Stenosis of nasal vestibule (Fig. 34.20)

- Nasal congestion
- Midfacial hypoplasia
- Otitis media
- Systemic side effects (47,48,49,50 and 51)
 - Infection
 - Aspiration
 - Increased intracranial pressure
 - Hypoxemia
 - Hypertension
 - o Apnea
 - Bradycardia and cardiac arrest
- Misplacements into esophagus or bronchus (52,53) (Figs. 34.21, 34.22 and 34.23)
 - Atelectasis
 - Pulmonary air leak
 - Loss of tube into esophagus
 - Across tracheoesophageal fistula
 - Displacement; accidental extubation (54)
- Obstruction (54)
- Kinking, proximally or distally
- Unrecognized disconnection from adapter or pressure source



FIG. 34.21. Radiograph demonstrating an endotracheal tube malpositioned in the bronchus intermedius, with resulting atelectasis of the right upper lobe and of the left lung. There is marked overaeration of the right middle and lower lobes but no pneumothorax as yet.

- Rupture of endotracheal tube (55)
- Foreign body from stylet left unrecognized in airway
- Swallowed laryngoscope light (56)
- Postextubation atelectasis (54)
- Increased airway resistance increasing work of breathing (57,58)



FIG. 34.22. Relatively uncommon malposition of an endotracheal tube in the left bronchus with atelectasis of much of the right lung.



FIG. 34.23. A: Radiograph suggesting that the endotracheal tube is in the right mainstem bronchus. Note the gaseous distension of the stomach and intestine. The wavy tube on the right is external. B: In the lateral view, the same endotracheal tube is easily seen to be in the esophagus (arrowheads) posterior to the trachea (arrows).

References

1. Todres ID, Crone RK. Experience with a modified laryngoscope in sick infants. Crit Care Med. 1981;9:544.

2. Yates AP, Harries AJ, Hatch DJ. Estimation of nasotracheal tube length in infants and children. Br J Anaesth. 1987;59:524.

R

3. De la Sierra Antona M, Lopez-Herce J, Ruperez M, et al. Estimation of the length of nasotracheal tube to be introduced in children. J Pediatr. 2002;140:772.

4. Freeman JA, Fredricks BJ, Best CJ. Evaluation of a new method for determining tracheal tube length in children. Anaesthesia. 1995;50:1050.

5. Shukla HK, Hendricks-Munoz KD, Atakent Y, et al. Rapid estimation of insertional length of endotracheal intubation in newborn infants. J Pediatr. 1997;131:561.

6. Kim KO, Um WS, Kim CS. Comparative evaluation of methods for ensuring the correct position of the tracheal tube in children undergoing open heart surgery. Anaesthesia. 2003;58:889.

7. Biskinis E, Herz M. Acquired palatal groove after prolonged orotracheal intubation. J Pediatr. 1978;92:512.

8. Moylan F, Seldin E, Shannon D, et al. Defective primary dentition in survivors of neonatal mechanical ventilation. J Pediatr. 1980;96:106.

9. Von Gonten AS, Meyer JB, Kim AK. Dental management of neonates requiring prolonged oral intubation. J Prosthodont. 1995;4:221.

10. Macey-Dare LV, Moles DR, Evans RD. Long-term effect of neonatal endotracheal intubation on palatal form and symmetry in 8–11 year-old children. Eur J Orthodont. 1999;21:703.

11. Lang M, Jonat S, Nikischin W. Detection and correction of endotracheal-tube position in premature neonates. Pediatr Pulmonol. 2002;34:455.

12. Woodall D, Whitfield J, Grunstein M. New recommendations for endotracheal tube positioning in the newborn infant [abstr]. Pediatr Res. 1981;15:733.

13. Wells TR, Wells AL, Galvis DA, et al. Diagnostic aspects and syndromal associations of short trachea with bronchial intubation. Am J Dis Child. 1990;144:1369.

14. Little LA, Koenig JC, Newth CJL. Factors affecting accidental extubations in neonatal and pediatric intensive care patients. Crit Care Med. 1990;18:163.

15. Jain A, Finer NN, Hilton S, Rich W. A randomized trial of suprasternal palpation to determine endotracheal tube position in neonates. Resuscitation. 2004;60:297.

16. Nicoll SJB, King CJ. Airway auscultation: a new method of confirming tracheal intubation. Anaesthesia. 1998;53:41. Cochrane Database Syst Rev.

17. Sutherland PD, Quinn M. Nellcor Stat Cap differentiates esophageal from tracheal intubation. Arch Dis Child Fetal Neonat Ed. 1995;73:184F.

18. Dejonge MH, White M. A comparison of two methods of oral endotracheal tube stabilization in neonatal patients. J Perinatol. 1998;18:463.

19. McMillan DD, Rademaker AW, Buchan KA, et al. Benefits of orotracheal and nasotracheal intubation in neonates requiring ventilatory assistance. Pediatrics. 1986;77:39.

20. Spence K, Barr P. Nasal versus oral intubation for mechanical ventilation of newborn infants. Cochrane Database Syst Rev. 2000;(2):CD000948.

21. Oei J, Hari R, Butha T, Lui K. Facilitation of neonatal nasotracheal intubation with premedication: a randomized controlled trial. J Paediatr Child Health. 2002;38:146.

22. Mayhall CG. The Trach Care® closed tracheal suctioning system: a new medical device to permit tracheal suctioning without interruption of ventilatory assistance. Infect Control Hosp Epidemiol. 1988;9:125.

23. Watkinson M, Rao JN. Endotracheal suction techniques in the neonate. Arch Dis Child. 1986;61:1147.24. Gould SJ, Howard S. The histopathology of the larynx in the neonate following endotracheal intubation. J Pathol. 1985; 146:301.

25. Brown MS. Prevention of accidental extubation in newborns. Am J Dis Child. 1988;142:1240.

26. Erenberg A, Nowak AJ. Appliance for stabilizing orogastric and orotracheal tubes in infants. Crit Care Med. 1984;12:669.

27. Richards S. A method for securing pediatric endotracheal tubes. Anesth Analg. 1981;60:224.

28. Markovitz BP, Randolph AG. Corticosteroids for the prevention and treatment of post-extubation stridor in neonates, children and adults. Cochrane Database Syst Rev. 2000;(2): CD001000.

29. Jansaithong J. The use of dexamethasone in the prevention of postextubation stridor in pediatric patients in PICU/NICU settings: an analytical review. J Soc Pediatr Nurs. 2001;6:182.

30. Mahieu HF, de Bree R, Ekkelkamp S, et al. Tracheal and laryngeal rupture in neonates: complication of delivery or of intubation?. Ann Otol Rhinol Laryngol. 2004;113:786.

31. Kolatat T, Aunganon K, Yosthiem P. Airway complications in neonates who received mechanical ventilation. J Med Assoc Thai. 2002;2:S455.

32. Pramanik AK, Sharma S, Wood BP. Traumatic hypopharyngeal pseudodiverticulum. Am J Dis Child. 1989;143:95.

33. Roberts D, McQuinn T, Beckerman RC. Neonatal arytenoid dislocation. Pediatrics. 1988;81:580.

34. Fan LL, Flynn JW, Pathak DR. Risk factors predicting laryngeal injury in intubated neonates. Crit Care Med. 1983;11:431.

35. Cotton RT. Prevention and management of laryngeal stenosis in infants and children. J Pediatr Surg. 1985;20:845.

36. Jones R, Bodnar A, Johnson D. Subglottic stenosis in newborn intensive care unit graduates. Am J Dis Child. 1981;135:367.

37. Sherman JM, Lowitt S, Stephenson C, et al. Factors influencing acquired subglottic stenosis in infants. J Pediatr. 1986; 109:322.

38. Couriel J, Phelan P. Subglottic cysts: a complication of neonatal endotracheal intubation? Pediatrics. 1981;68:103.

39. Angelos GM, Smith DR, Jorgenson R, et al. Oral complications associated with neonatal oral tracheal intubation: a critical review. Pediatr Dent. 1989;11:133.

40. Molteni RA, Bumstead DH. Development and severity of palatal grooves in orally intubated newborns. Am J Dis Child. 1986;140:357.

41. Seow WK, Humphrys C, Tudehope DE. Increased prevalence of developmental dental defects in low birth weight, prematurely born children: a controlled study. Pediatr Dent. 1987; 9:221.

42. Kopra DE, Creighton PR, Buckwald S, et al. The oral effects of neonatal intubation. J Dent Res. 1988;67:165.

43. Kahn DJ, Spinazzola R. Acquired oral commissure defect: a complication of prolonged endotracheal intubation. J Perinatol. 2005;25:612.

44. Gowdar K, Bull M, Schreiner R, et al. Nasal deformities in neonates. Their occurrence in those treated with nasal continuous positive airway pressure and nasal endotracheal tubes. Am J Dis Child. 1980;134:954.
45. Rotschild A, Dison PJ, Chitayat D, et al. Midfacial hypoplasia associated with long-term intubation for bronchopulmonary dysplasia. Am J Dis Child. 1990;144:1302.

46. Halac E, Indiveri DR, Obergaon RJ, et al. Complication of nasal endotracheal intubation. J Pediatr. 1983;103:166.

47. Storm W. Transient bacteremia following endotracheal suctioning in ventilated newborns. Pediatrics. 1980;65:487.

48. De Dooy J, Leven M, Stevens W, et al. Endotracheal colonization at birth is associated with a pathogendependent pro- and antiinflammatory cytokine response in ventilated preterm infants: a prospective cohort study. Pediatr Res. 2004;56:547.

49. Marshall TA, Deeder R, Pai S, et al. Physiologic changes associated with endotracheal intubation in preterm infants. Crit Care Med. 1984;12:501.

50. Goitein KJ, Rein AJ, Gornstein A. Incidence of aspiration in endotracheally intubated infants and children. Crit Care Med. 1984;12:19.

51. Frieson RH, Honda AT, Thieme RE. Changes in anterior fontanel pressure in preterm neonates during tracheal intubation. Anesth Analg. 1987;66:874.

52. Bagshaw O, Gillis J, Schell D. Delayed recognition of esophageal intubation in a neonate: role of radiologic diagnosis. Crit Care Med. 1994;22:2020.

53. Buchino JJ, Keenan WJ, Pletsch JB, et al. Malpositioning of the endotracheal tube in infants with tracheoesophageal fistula. J Pediatr. 1986;109:524.

54. Rivera R, Tibballs J. Complications of endotracheal intubation and mechanical ventilation in infants and children. Crit Care Med. 1992;20:193.

55. Spear RM, Sauder RA, Nichols DG. Endotracheal tube rupture, accidental extubation, and tracheal avulsion: three airway catastrophes associated with significant decrease in peak pressure. Crit Care Med. 1989;17:701.

56. Naumovski L, Schaffer K, Fleisher B. Ingestion of a laryngoscope light bulb during delivery room resuscitation. Pediatrics. 1991;87:581.

57. Lesouef PN, England SJ, Bryan AC. Total resistance of the respiratory system in preterm infants with and without endotracheal tube. J Pediatr. 1984;104:18.

58. Perez Fontan JJ, Heldt GP, Gregory GA. Resistance and inertia of endotracheal tubes used in infants during periodic flow. Crit Care Med. 1985;13:1052.

35-Tracheotomy

Hosai Hesham Gregory J. Milmoe

A. Indications

- Prolonged need for ventilator support â€"most common
- Acquired subglottic stenosis after prolonged intubation
- Craniofacial abnormalities with severe airway obstruction, e.g., Pierre-Robin sequence, Pfeiffer syndrome, Treacher Collins syndrome
- Congenital bilateral vocal cord paralysis
- Laryngeal web, subglottic hemangioma
- Congenital tracheal stenosis, severe tracheomalacia
- Congenital neuromuscular disease with insufficient respiratory effort
- Neurologic disease with aspiration risk, central apnea, or intractable seizures

B. Contraindications

- Unstable physiology wait until stabilized
 - Sepsis
 - Pneumonia not yet controlled
 - Pulmonary instability requiring high inspiratory pressures (PIP >35-40) or need for high-frequency ventilation
 - o Cardiovascular instability, e.g., shunting, arrhythmia, or hypotension
 - Evolving renal or neurologic injuries
- Distal obstruction not relieved by tracheostomy
 - Congenital stenosis at the carina
 - External compression from mediastinal mass
- Congenital anomalies that make the trachea relatively inaccessible
 - Massive cervical hemangioma bleeding issues
 - Massive cervical lymphangioma severe distortion of neck anatomy
 - Massive goiter might be manageable medically
 - Chest syndromes with severe kyphoscoliosis or tracheal distortion

C. Precautions

• Patient should be stable (see Contraindications); anticipate need for increased pulmonary support temporarily to counter atelectasis and reactive secretions from surgical stimulation.

- Tracheotomy tubes allow for air leak through stoma and through larynx. In contrast, an endotracheal tube fits more snugly at the cricoid, creating a more closed system for ventilation.
- Neonates are less able to tolerate bacteremia, so use perioperative antibiotic to cover skin flora.
- If the patient is not currently intubated, have endoscopy equipment available and discuss intubation options with the anesthesiologist.
- The infant larynx differs from that of the adult and older child (Fig. 35.1).
 - More pliable and mobile
 - Relatively higher in neck
 - Thymus and innominate artery can override trachea in surgical field
- This procedure should be done only in a facility where there is appropriate support for postoperative management.

D. Equipment

- Prep tray with brushes, towels, and betadine
- Tracheotomy tray
 - Scalpel with no. 15 blade
 - Hemostats
 - Small scissors (iris, tenotomy, small Mayo)
 - Retractors Senn or Ragnel
 - Suction no. 7 Frazier
 - Forceps Adson
- Sutures: 3-0 and 4-0 nonabsorbable on small, curved needles
- Neonatal tracheotomy tubes
 - Have several calibers available
 - Standard tubes are noncuffed, but in special circumstances a cuff may be needed.

E. Technique

- Check instruments, sutures. and available tracheotomy tubes.
- On patient arrival in operating room, apply monitors, check intravenous line, and confirm satisfactory ventilation through endotracheal tube.
- Anesthesia team proceeds with inhalation agents, oxygen supplementation, and intravenous agents as needed for satisfactory level of general anesthesia.
- Position patient with neck extended using shoulder roll.
- Remove nasogastric tube to avoid confusion when palpating trachea. Do not place esophageal stethescope either.
- Inject skin incision and the deeper tissues with local anesthetic (0.5 to 1.0 mL of Â¹/₂% lidocaine with 1:200,000 epinephrine).
- Prep the surgical site from above the chin to below the clavicles. Give IV antibiotic to cover skin flora.
- Drape the patient with surgical towels, allowing the anesthesiologist access to the endotracheal tube and the securing tape.
- Identify the following landmarks: suprasternal notch, chin, midline, trachea, and cricoid. In small neonates, the cricoid may be difficult to palpate.
- The skin incision is made roughly midway between the sternal notch and the cricoid, either vertically or horizontally. Both tend to heal as a circular stoma, but the horizontal has a slightly better cosmetic effect, while the vertical allows more exposure in the midline.
- Excess subcutaneous fat can be excised with cautery.
- Identify the strap muscles and repeatedly palpate the trachea to confirm the midline. Split the raphe to separate the muscles.

- Grab the fascia of the strap muscles with hemostats to retract them outward and laterally, thereby exposing the thyroid gland, cricoid, and trachea.
- Senn retractors can then be placed on either side of the trachea for better view.
- The thyroid gland can usually be displaced by blunt dissection to expose the tracheal rings. If not, then the isthmus is divided and suture ligated.
- Place vertical stay sutures in paramedian position at the level where tracheal entry is planned usually the third and fourth ring (Fig. 35.2).
- Incise trachea vertically for 2 or 3 rings, depending on the size needed for the tube employed.
- The anesthesiologist can then loosen the tape and withdraw the endotracheal tube until the tip is just visible (Fig. 35.3).
- The appropriate tracheostomy tube is then placed with the flange parallel to the trachea so that the tube more easily enters the trachea and passes posteriorly before rotating the flange 90 degrees back to its routine orientation.
- The anesthesiologist confirms placement by checking end-tidal carbon dioxide and oxygen saturation, as well as auscultation of both sides.



FIG. 35.1. Sagittal section. Larynx lies more cephalad than in adult. Note the proximity of the thyroid isthmus to the tracheal rings. (Drawing contributed by John Bosma, M.D.)



FIG. 35.2. Placement of stay sutures through the tracheal wall.

- The tracheostomy tube is then secured with twill tape tied around the neck firmly. Once tied, only one finger should fit between the tape and the neck when the baby's neck is in neutral position.
- The stay sutures are then secured to the chest with tape labeled as to correct side (Fig. 35.4).
- Transport the patient back to intensive care with a backup endotracheal tube and laryngoscope.
- Obtain chest radiograph on arrival in unit to check tube position and lung status.



FIG. 35.3. Artistic conception of view through tracheal incision with the tip of the endotracheal tube visible. Stay sutures hold cartilages open.

F. Postoperative Management

- Provide intensive nursing (see Precautions).
- Keep spare tracheostomy tubes at bedside (same size and one smaller).
- Replace nasogastric tube for nutrition and to avoid aerophagia.
- Suction secretions as needed to avoid plugging. For first 24 hours, be liberal with saline irrigation.
- Make sure ventilator tubing is not pulling on tracheostomy tube.
- Be aggressive with wound care so that stoma heals quickly and thereby limits granulation. Clean once a shift with half-strength peroxide and cotton swabs, then apply antibiotic ointment.
- First tracheostomy change is done by surgical team in 4 to 7 days. Thereafter weekly changes are sufficient.



FIG. 35.4. Fixation of stay sutures. As soon as the position of the tracheostomy tube is confirmed and stomal ventilation is started, the tube may be fixed. Equal tension is kept on the stay sutures during taping. Right suture is marked to avoid confusion in future placement.

G. Early Complications (0 to 7 days)

- Bleeding: thyroid, venous, arterial
- Accidental decannulation or displacement in neck â€"stay sutures are the child's lifeline back to the trachea to allow replacement of the tube.
- Plugging of tube with secretions (Fig. 35.5)
 - Avoid by increasing humidity, saline irrigation, and suctioning.
- Infection of wound or pneumoniaâ€"avoid by local care, and by taking care of secretions.
- Air leaks
 - Pneumothorax may need chest tube
 - Pneumomediastinum serial films
 - Subcutaneous emphysema usually limited (avoid occlusive dressing)
- Tracheoesophageal fistula iatrogenic

H. Late Complications (after 1 week)

- Obstruction and decannulation remain ongoing risks that require vigilant care.
- Stomal infection and granulation â€"avoided by careful wound care
- Proximal tracheal granuloma â€" commonly occurs at the point where the tube rubs against the superior aspect of the tracheal opening, creating an obstruction between the vocal cords and the tube that can impede routine tracheostomy tube changes. This requires operative removal.
- Distal tracheal granulation â€"from overly aggressive suctioning or tube angulation causing rubbing of the tip against the tracheal wall. Hallmark sign is bloody secretions.



FIG. 35.5. Total obstructions of tracheostomy tubes. A: Mucus plug incompletely suctioned. B: Dry mucus plug pushed deeper by a suction catheter.

- Stenosis preventing decannulation later on
 - Part of original pathology requiring tracheostomy
 - Ongoing obliteration from active inflammatory factors
 - Consequence of procedure itself from stomal collapse or distal cicatrix

• Tracheocutaneous fistula after tube removalâ€"normal physiologic sequela, but needs secondary procedure for closure

References

1. Sisk EA, Kim TB, Schumacher R, et al. Tracheotomy in very low birth weight neonates: indications and outcomes. Laryngoscope. 2006;116:928–933.

2. Wooten CT, French LC, Thomas RG, et al. Tracheotomy in the first year of life: outcomes in term infants, the Vanderbilt experience. Otolaryngol Head Neck Surg. 2004;134:365–369.

3. Kremer B, Botos-Kremer AI, Eckel HE, et al. Indications, complications and surgical techniques for pediatric tracheostomiesâ€"an update. J Pediatr Surg. 2002;37:1556â€"1562.

4. Crysdale WS, Feldman RI, Naito K. Tracheostomies: a 10 year experience in 319 children. Ann Oto Laryngol. 1988;97:439-443.

36-Thoracostomy Tubes

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Thoracostomy Tubes

Pulmonary air leak is an anticipated risk of mechanical ventilation. Drainage of air and/or fluid accumulation in the chest is an important and necessary skill and is often performed on an emergency basis. Thoracostomy tubes are used in neonatal intensive care units for evacuation of air or fluid from the pleural space. The procedure is often performed because of an emergency. In addition to recognizing pathologic states that necessitate chest tube insertion, intensive care specialists are frequently involved in placement, maintenance, troubleshooting, and discontinuation of chest tubes. As with any surgical procedure, complications may arise. Appropriate training and competence in the procedure may reduce the incidence of complications. This chapter reviews current indications for chest tube placement, insertion techniques, and equipment. Guidelines for chest tube maintenance and discontinuation are also discussed. A. Indications

- Evacuation of pneumothorax
 - Tension
 - Lung collapse with ventilation/perfusion abnormality
 - Bronchopleural fistula
- Evacuation of large pleural fluid collections
 - Significant pleural effusion
 - Postoperative hemothorax
 - Empyema
 - Chylothorax
 - Extravasated fluid from a central venous line
- Extrapleural drainage after surgical repair of esophageal atresia and/or tracheoesophageal fistula
- B. Contraindications
 - Small air or fluid collection without significant hemodynamic symptoms

• Spontaneous pneumothorax that, in the absence of lung disease, is likely to resolve without intervention

C. Equipment

Sterile

- General all-purpose tray with no. 15 surgical blade and curved hemostats
- Gloves
- Surgical drapes
- Transillumination device
- Thoracostomy tube: Techniques of insertion differ with each type. See original references for description of technique variations (1,2 and 3).
 - Polyvinyl chloride (PVC) chest tube with or without trocar, in sizes 8, 10, and 12 French (Fr)
 - Pigtail catheter for pleural drainage (Fig. 36.1)
 - PVC with pigtail at 90-degree angle to shaft (1)
 - 8 to 10 Fr
 - Total length 10 cm
 - Insertion with or without trocar
 - Polyurethane modified vascular catheter with pigtail in same plane as shaft (2)
 - 8.5 Fr
 - Total length 15 cm
 - Insertion guide wire and dilator for insertion by Seldinger technique
 - Cook catheter (C-PPD-500/600-MP8561; Cook, Bloomington, IN, USA) (3)
 - 5 and 6 Fr
 - Cutting needle tip joined to a biopsy needle shaft with a collar that prevents the catheter from sliding up the needle during insertion
- Evacuation device
 - Infant thoracostomy tube set: Several commercially available units are appropriate for infants (Fig. 36.2).
 - Evacuation rate (4)
 - With single tube, capacity depends on level of water in chamber (cm H₂O).



FIG. 36.1. Pigtail catheter for pleural drainage (Fuhrman pleural drainage set). (Illustration provided by Cook Critical Care, Bloomington, IN, USA).

- With multiple tubes, capacity also depends on applied vacuum.
- Negative pressure of 20 cm H₂O evacuates more than 4 L of air/min in experimental setting (4).
 - Appropriate starting point for most infants with lung disease on ventilators is 10 to 15 cm H₂O
 - Potentially inadequate in a case of bronchopleural fistula
 - Excessive suction pressure may draw tissue into the side holes of the chest tube and could also be potentially harmful in changing intrapulmonary air flow in presence of smaller pleural leak (always start with 10 cm H₂O).

Measured rates across bronchopleural fistulas in infants have indicated ranges from 30 to 600 mL/min (5). If suction pressure is too high, gas flow to alveoli may be diverted across a fistula. The pressure and flow applied to the endotracheal tube also directly influence flow across a fistula (5). Because there are many interactive factors influencing how much air might have to be evacuated, there can be no single best suction level for all patients; the most effective, least harmful level has to be determined for each situation (6).

- Nonabsorbable suture on small cutting needle, 4.0
- Cotton-tipped applicators
- Semipermeable transparent dressing
- Antibiotic ointment
- Petroleum gauze

Nonsterile

- Tincture of benzoin
- Â¹/₂-in adhesive tape
- Towel roll

D. Factors Influencing Efficiency of Air Evacuation

- Contiguity of air to chest tube portals; they must be patent
 - In supine infant, air accumulates in the medial, anterior, or inferior hemithorax, making low anterior location for tip of tube ideal for evacuation (7).



FIG. 36.2. One model of an underwater drainage system, demonstrating the three necessary chambers. Systems now are compact and easy to set up and read. This system is set at 22 cm H_2O , which would be necessary only for a rapid rate of air accumulation.

- \circ Negative pressure on chest tube may draw tissue into side portals and occlude them.
- Rate of air accumulation proportional to
 - Airway flow and pressure

Dennis et al. (8) demonstrated in experimental rabbit models that a positive end-expiratory pressure level >6 cm H_2O resulted in greater air leak than peak inspiratory pressure, up to 30 cm H_2O .

- \circ Size of fistula or tear
- Infant position

If the affected side is the dependent hemithorax and therefore is splinted, there is a lower rate of air leak than if the affected side is not dependent or is elevated (9).

- Rate of evacuation
 - Directly proportional to
 - Internal radius of chest tube (r⁴)
 - Pressure gradient across tube (DP)
 - Suction pressure applied

The negative pressure applied may affect intrapleural pressure only in the immediate vicinity of the tip of the tube (4).

- Positive intrathoracic pressure during exhalation, spontaneous or mechanical
- o Inversely proportional to length of tube and viscosity

Poiseuille's law regarding flow across a tube is $F = DPI \in r^4/8hl$, where F = flow; DP = pressure gradient; r = radius; h = viscosity; and l = length.

E. Precautions

- Anticipate which infant is at risk of developing pulmonary air leakage and keep equipment for diagnosis and emergency evacuation at hand (6,10,11).
- Recognize that transillumination may be misleading (12,13).
 - True positive
 - Follows shape of thoracic cavity (not corona of light source)
 - Varies with respiration and position
 - Has larger area compared with corona of light at another normal site
 - False positive
 - Subcutaneous edema
 - Subcutaneous air
 - Severe pulmonary interstitial emphysema
 - False negative
 - Thick chest wall
 - Darkly pigmented skin
 - Area over air accumulation obscured by dressing/monitor probe
 - Weak light because of fiber-optic deterioration or voltage turned too low
 - Room too light
 - Abnormal color vision in observer
- Distinguish pleural air collections from skin folds, thymus, Mach effect, artifacts, or other nonpleural intrathoracic air collections on radiograph (Figs. 36.3,36.4,36.5 and 36.6) (7,14).
- Select the appropriate insertion site (Figs. 36.7 and 36.8).

Allen et al. (15) recommend inserting the thoracostomy tube in the anterosuperior portion of the chest wall, in the first to third intercostal space at the midclavicular line, to ensure anterior placement of the chest tube tip. However, although an anterior insertion may be appropriate for the right-angled
pigtail tube used by Allen et al., a properly placed lateral tube will have its tip anterior but, more important, will not leave a (more visible) scar on the anterior chest and completely avoids the nipple (see Fig. 36.19).



FIG. 36.3. Sequential radiographs. A: Anteroposterior radiograph demonstrating a cystic lucency at the left base behind the heart (arrows) that resembles the artifact caused by taking a film through the hole in the top of an incubator. Note also the coarse, irregular lucencies of interstitial emphysema (PIE) in the left lung. B: Lateral film showing the lucency to be real (arrows) and, in this case, a pneumomediastinum located most probably in the left inferior pulmonary ligament. C: PIE and air in the pulmonary ligament are often harbingers of impending pneumothorax, in this case, a tension pneumothorax. Note low position of endotracheal tube.



FIG. 36.4. Radiographic artifact of cystic lucency behind the heart (arrows) caused by taking film through top of incubator. The lateral film was negative, therefore excluding a cystic pulmonary lesion or air in the pulmonary ligament.

- Reduces complications
- Facilitates insertion of thoracostomy tube into appropriate position
 - Anteromedial tip for air collections
 - Posterior tip for fluid accumulation (Fig. 36.9A, B).
- While inserting the chest tube, allow some air to remain within pleural space as protective buffer between lung and chest wall (6).
 - Use emergency pneumothorax evacuation only if patient is critically compromised. If emergency evacuation is used, remove air only until vital signs are stable.
 - Position infant so point of entry is the most elevated area of the chest.
 - Allows air to rise to provide protective buffer.
 - Direct tip of the chest tube anteriorly toward the apex of the thorax.
- Consider the possibility that a rapid, complete evacuation may cause an abrupt increase in mean arterial blood pressure and cerebral blood velocity to undesirable, supranormal levels (16).
- Avoid positioning infant in lateral decubitus position for any longer than necessary with "normal†lung dependence, thereby further compromising ventilation.
- To prevent laceration of lung parenchyma, avoid inserting needles beyond parietal pleura for diagnostic or emergency taps. Use a straight clamp perpendicular to the needle shaft to limit depth of penetration (Fig. 36.10).
- Do not use purse-string suturing of the incision site, because resulting scars tend to pucker (6,17) (see Fig. 36.19).
- Recognize that air leaks are likely to persist after initial evacuation in the presence of continuing lung disease or positive-pressure ventilation. Air leaks resolve in 50% of patients within the first 4 days after chest tube placement, and 83% resolve after 7 days (18).
 - Continue to watch for patency of the chest tube (Fig. 36.11).
 - Verify the correct position of the tube.

- Modify positive-pressure ventilator patterns to minimize risk of further air leaks (10).
 - Decrease inspiratory time.
 - Decrease mean airway pressure.
- Position infant with the side of the pleural gas leak dependent (9).

F. Technique (See also Procedures DVD)

Insertion of Anterior Tube for Pneumothorax

- Determine location of air collection.
 - Physical examination

Auscultation of the small neonatal chest may be misleading because the breath sounds normally are bronchotubular and may be relatively well transmitted across an air-filled hemithorax. In addition, shift of the point of maximal impulse toward the other side is unusual in the presence of noncompliant lungs. Physical findings of acute abdominal distention, irritability, and cyanosis or a change in transthoracic impedance suggest an air leak but not its location (19,20). Supplementary diagnostic procedures are usually necessary.

- Transillumination (12)
- Radiograph (7,21)
- Support the infant with artificial ventilation as required. The majority of infants with a pneumothorax requiring a chest tube also need ventilatory support.
- Monitor vital signs. Move any electrodes from the operative site to alternative monitoring areas.
- Position infant with affected side elevated 60 to 75 degrees off the bed and support the back with a towel roll. Secure arm across the head, with shoulder internally rotated and extended (Fig. 36.12A).

This position is very important because it allows air to rise to the point of tube entry into the thoracic cavity, outlines the latissimus dorsi muscle, and encourages the correct anterior direction of the tube.

- Prepare the skin with an iodophor antiseptic over the entire lateral portion of chest to the midclavicular line. Blot excess antiseptic and allow skin to dry.
- •



FIG. 36.5. A: Anteroposterior radiograph demonstrates ventral air over the hemidiaphragms and around the heart (arrowheads). The sometimes difficult question of pneumothorax versus pneumomediastinum is answered by the decubitus films. B: The left lateral decubitus radiograph (right side up) shows that the right-sided gas is a pneumothorax (arrowheads). C: The right decubitus film indicates that the adventitial air fails to come up over the lung and is located in the mediastinum (arrowheads). This important distinction is made obvious by the decubitus radiographs.

- Drape surgical area from third to eighth ribs and from latissimus dorsi muscle to midclavicular line (Fig. 36.12B). Using a transparent aperture drape allows continued visualization of landmarks.
- Locate essential landmarks (Fig. 36.12C).
 - Nipple and fifth intercostal spaces
 - Midaxillary line
 - Skin incision site is at point midway between midaxillary and anterior axillary lines in the fourth or fifth intercostal space. A horizontal line from the nipple is a good landmark for identifying the fourth intercostal space. Keep well away from breast tissue (22).
- Remove trocar from tube.



FIG. 36.6. A: On this anteroposterior supine film, there is a line that parallels the chest wall (arrowheads), which suggests the presence of a pneumothorax. B: This left decubitus film (right side up) confirms this line to be a skin fold, negative for air. When there is a question of potential adventitial air or of the anatomic location of real adventitial air, a decubitus film with the side in question up is the most important radiographic study.

We do not recommend using a trocar during tube insertion because of the greater likelihood for lung perforation. Dissection to the pleura should be performed as described here, with puncture of the pleura by the tip of the closed forceps, not by a trocar. Should a trocar be used after dissecting to the pleura, there should be a straight clamp perpendicular to the shaft at 1 to 1.5 cm from its tip to avoid penetrating too deeply (Fig. 36.10).

- Estimate length for insertion intrathoracic portion of tube (skin incision site to mid-clavicle). This should be approximately 2 to 3 cm in a small preterm infant and 3 to 4 cm in a term infant. (These are approximate guidelines only.)
- Infiltrate skin at incision site with 0.125 to 0.25 mL of 1% lidocaine.
- Using a no. 15 blade, make incision through skin the same length as chest tube diameter or no more than 0.5 to 1.0 cm (Fig. 36.12C).
- Puncture pleura immediately above the fifth rib by applying pressure with index finger (Fig. 36.12D).
 - Place the forefinger as shown in Fig 36.12D and not further forward on the forceps, to prevent the tip from plunging too deeply into the pleural space.
 - A definite give will be felt as the forceps tip penetrates the pleura; there may also be an audible rush of air.



FIG. 36.7. Anterior versus posterior position of the tube for drainage of air or fluid. Because air collects anteromedially in the supine neonate, the posterior tip is less appropriate.

- After puncturing pleura, open hemostat just wide enough to admit chest tube.
- Leaving hemostat in place, thread tube between opened tips to the predetermined depth (Fig. 36.12E).
 - Alternatively, insert closed tips of mosquito hemostat into side port of tube to its end. The disadvantage of this method is that the forceps will have to be withdrawn from the opening in the chest; it is common that the intercostal muscles then render the opening undetectable (22,23).



FIG. 36.8. Sequential radiographs in patient with right pneumothorax. An air collection in supine neonates (A) is most effectively treated with an anteromedial chest tube (B, C). The medial extension is falsely exaggerated by the slight right posterior oblique position of the chest. Pulling this tube back might put the side holes outside the pleural space. There is a

pneumomediastinum, most evident on the lateral view, not drained by the pleural tube. Note the nuchal air on all three films.

- Direct chest tube cephalad, toward apex of thorax (midclavicle), and advance tip to midclavicular line, ensuring that all side holes are within pleural space.
- \circ $\;$ Observe for humidity or bubbling in chest tube to verify intrapleural location.
- Connect tube to vacuum drainage system and observe fluctuations of meniscus and pattern of bubbling (Fig. 36.11). Avoid putting tension on tube.
- Secure chest tube to skin with suture (Fig. 36.13A).



FIG. 36.9. A: Photograph of a pigtail catheter placed posteriorly for pleural fluid drainage. B: Serosanguineous pleural fluid collection into the chest tube set.

• Use one suture to close end of skin incision and make airtight seal with chest tube. Tie ends of suture around tube in alternating directions, without constricting tube.

Because using a traditional purse-string suture to secure the tube leaves an unsightly scar, we do not recommend it. Unless skin incision has been made unnecessarily long, a single suture is usually sufficient.



FIG. 36.10. Chest wall in cross section. If there is need to use a needle or trocar to enter the pleural space, its depth of penetration should be limited by a perpendicular clamp.

- Apply tincture of benzoin to chest tube near chest wall and to skin several centimeters below incision. When tacky, encircle tube with a 2-in length of tape, leaving tab posterior (Fig. 36.13B).
- Place suture through skin and tab of tape to stabilize chest tube in straight position (Fig. 36.13B).
- Alternatively, secure tube with tape bridge (Fig. 36.14) or clear adhesive dressing (the latter may not be optimal; chest tubes tend to function optimally when allowed to exit from the skin at as close to a 90-degee angle as possible).
- Apply antibiotic ointment or petroleum gauze around skin incision. Cover with small semiporous transparent dressing.

It is important not to cover the wound with a heavy dressing, which restricts chest wall movement, obscures tube position, and makes transillumination more difficult. If the position of tube is in doubt, secure with temporary tape bridge before covering with dressing until correct position is confirmed.

• Verify proper positioning of tube.



FIG. 36.11. Evaluation of a chest tube: Flow chart to determine how well a chest tube is evacuating pleural air leak and when tube should be removed.

• Anteroposterior and lateral radiographs (6,24,25 and 26)

Both views are recommended to detect anterior course of tube. See Tables 36.1 and 36.2 for radiographic clues on malpositions. Malpositioned tube tip results in an increased risk of complications and/or poor air evacuation. Chest radiograph should confirm that the side holes are within the chest cavity.

TABLE 36.1 Clues to Recognize Thoracostomy Perforation of the Lung

- 1. Bleeding from endotracheal tube
- 2. Continuous bubbling in underwater seal
- 3. Hemothorax
- 4. Blood return from chest tube
- 5. Increased density around tip of tube on radiograph
- 6. Persistent pneumothorax despite satisfactory position on frontal view
 - Tube lying neither anterior nor posterior to lung on lateral view
- 8. Tube positioned in fissure
- Pattern of bubbling (Fig. 36.11)

7.

• Strip tube if meniscus stops fluctuating or as air evacuation decreases. Take extreme care not to dislodge tube by holding tube firmly with one hand close to chest wall.

Insertion of Posterior Tube for Fluid Accumulation

The technique is similar to that for an anteriorly positioned tube, with the following differences.

- Position infant supine, elevating the affected side by 15 to 30 degrees from the table. Secure the arm over the head (Fig. 36.15).
- Prepare skin over lateral portion of hemithorax from anterior to posterior axillary line.

TABLE 36.2 Clues to Thoracostomy Tube Positioned in Fissure

- 1. Major interlobar fissure
 - a. Frontal view: Upper medial hemithorax
 - b. Lateral view: Oblique course posterior and upward
 - Minor fissure (on right)
 - a. Horizontal course toward medial side of lung
- Make skin incision of 0.5 to 0.75 cm in length, just behind the anterior axillary line in the fourth to sixth intercostal space and following direction of rib.
 - Fourth or fifth space for high posterior tube tip
 - Sixth space for low posterior tube tip
- Taking care to position forceps tip immediately above a rib to avoid the intercostal vessels that run under the inferior surface, penetrate the pleura as described for anterior chest tube.
- Insert tube only deeply enough to place side holes within pleural space.
- Collect drainage material for culture, chemical analysis, and volume.
- Connect to underwater seal drainage system that includes a specimen trap.
- Strip tube regularly.

2.

• Monitor and correct any imbalance caused by loss of fluid, electrolytes, protein, fats, or lymphocytes.

Removal of Thoracostomy Tube

- Ascertain that tube is no longer functioning or needed.
 - Evaluate as suggested in Fig. 36.11.
 - Leave chest tube connected to water seal without suction for 2 to 12 hours. Do not clamp tube.
 - Transilluminate to detect reaccumulation.
 - Obtain radiograph.
 - Document absence of significant drainage.
- Assemble equipment.
 - Antiseptic solution
 - Sterile gloves
 - Scissors
 - Forceps
 - Petroleum gauze cut and compressed to 2-cm diameter
 - Gauze pads, 2 x 2 in
 - 1-in tape
- Cleanse skin in area of chest tube with antiseptic.
- Release tape and suture holding tube in place. Leave wound suture intact if skin is not inflamed.
- Palpate pleural entry site and hold finger over it to prevent air from entering chest as tube is withdrawn until gauze is applied. After removing tube, approximate wound edges and place petroleum gauze over incision. Keep pressure on pleural wound until dressing is in place.

- Cover petroleum gauze with dry, sterile gauze. Limit taping to as small an area as possible so that transillumination will be possible.
- Remove sutures when healing is complete.
- G. Complications
 - Misdiagnosis with inappropriate placement
 - Burn from transillumination devices (27)



FIG. 36.12. Insertion of a soft chest tube. A: Position the infant with back support so the point of tube entry will be highest. Fix arm over the head without externally rotating it. Note the midaxillary (MA) line and the line from the nipple through the fourth intercostal space (ICS). B: Drape so head of the infant is visible. C: Same landmarks without the drape, showing the incision in the fourth ICS in the MA line with entry into the chest at the intersection of the nipple line and the MA line. D: Turning the hemostat to puncture into the pleura in the fourth ICS. E: With the index finger marking the fourth ICS puncture site, the tube may now be passed between the hemostat blades, along the tunnel into the pleural space.

- Trauma
 - Lung laceration or perforation (28,29) (Fig. 36.16)
 - Perforation and hemorrhage from a major vessel (axillary, pulmonary, intercostal, internal mammary) (15,30) (Fig. 36.17)
 - Puncture of viscus within path of tube (Fig. 36.18)
 - Residual scarring (17) (Fig. 36.19)
 - Permanent damage to breast tissue (17)
 - Chylothorax (31)
- Nerve damage
 - Horner syndrome caused by pressure from tip of right-sided, posterior chest tube near second thoracic ganglion at first thoracic intervertebral space (32)
 - Diaphragmatic paralysis or eventration from phrenic nerve injury (33,34 and 35)
- Misplacement of tube
 - Tube outside pleural cavity in subcutaneous placement (Fig. 36.20)
 - Side hole outside pleural space (Fig. 36.21)
 - Tip across anterior mediastinum (Fig. 36.22)
- Equipment malfunction
 - Blockage of tube by proteinaceous or hemorrhagic material
 - \circ $\;$ Leak in evacuation system, usually at connection sites
 - Inappropriate suction pressures (36,37) (Fig. 36.11)
 - Excessive pressure
 - Aggravation of leak across bronchopleural fistula
 - Interference with gas exchange
 - Suction of lung parenchyma against holes of tube
 - Inadequate pressure with reaccumulation
- Infection
 - \circ Cellulitis
 - Inoculation of pleura with skin organisms including Candida (38)
- Subcutaneous emphysema secondary to leak of tension pneumothorax through pleural opening



FIG. 36.13. Securing a chest tube. A: Make the incision site airtight with the tube. Do not use a purse-string suture around the incision, because it will form a puckered scar. The initial incision should be made small enough to require only a single suture. B: After painting the tube and skin with benzoin, encircle the suture around the tube or attach a bandage and suture it to the skin.



FIG. 36.14. Tape bridge. A: Two tape towers. B, C: Bridge under the tube and towers overlapping on top. D: Additional cross tape to keep the chest tube flat without kinking.



FIG. 36.15. Insertion of a posterior chest tube. With the infant supine, the incision is in or just below the anterior axillary line with the tube entry into the pleura more posteriorly Take care to enter pleural space over the top of a rib.



FIG. 36.16. Postmortem examination of infants who died with uncontrolled air leaks. A: Perforation of the right superior lobe by a chest tube inserted without a trocar, demonstrating that virtually any tube can penetrate into the lung. B: Perforation of the left upper lobe by a chest tube (arrow).



FIG. 36.17. Posterior view of thoracic organs. Traumatic hemorrhage of the left upper lobe was due to perforation by a thoracostomy tube.



A FIG. 36.18. Postmortem examination of an infant with bilateral pneumothorax, pneumomediastinum, and pneumoperitoneum secondary to pulmonary air leaks. Attempted needle aspirations, as seen by multiple skin puncture sites of the pneumomediastinum and pneumothorax (A), resulted in needle punctures of the liver (arrows, B) with peritoneal hemorrhage.



FIG. 36.19. Scar from thoracostomy insertion, emphasizing the importance of avoiding the breast area. Massage of a healed wound with cocoa butter helps break down adhesions that lead to dimpling at the scar.



FIG. 36.20. The thoracostomy tube is completely outside the pleural space on this slightly oblique chest film. Note that the long feeding tube is not in an appropriate position for transpyloric feeding. Indwelling tubes may dislodge when other emergency procedures are performed.



FIG. 36.21. The side holes of both thoracostomy tubes are outside the pleural space on this radiograph.



FIG. 36.22. The tip of the thoracostomy tube has been advanced too far medially and is kinked against the mediastinum. Withdrawing the tube 1 or 2 cm would improve drainage at the medial pneumothorax. Note the endotracheal tube tip in the right mainstem bronchus.



FIG. 36.23. Emergency evacuation with a vascular cannula. Puncture the skin and enter the pleura at a 45-degree angle immediately above a rib.

- Aortic obstruction with posterior tube (39)
- Loss of contents of pleural fluid
 - Water, electrolytes, and protein (effusion)
 - Lymphocytes and chylomicrons (chylothorax)

Emergency Evacuation of Air Leaks

Life-threatening air accumulations require emergency evacuation. This provides temporary relief to the patient while preparing for thoracostomy tube placement. The following techniques using modified equipment are less traumatic than using straight needles or scalp vein sets. We suggest using an anterior approach for emergency evacuation because position will not interfere with the preparation of the lateral chest site for an indwelling chest tube.

Tubes used for emergency evacuation require suction pressures as high as 30 to 60 cm H_2O to overcome the resistance of their small diameters (40). This requirement and their tendency to occlude make these cannulas unreliable for continuous drainage of a significant air leak.

A. Indications

Temporary evacuation of life-threatening air accumulations while preparing for permanent tube placement

B. Contraindications

- When patient's vital signs are stable enough to allow placement of permanent thoracostomy tube without prior emergency evacuation
- When air collection is likely to resolve spontaneously without patient compromise (nontension pneumothorax).

Use of Angiocatheter (41)

A. Equipment

- Sterile gloves
- Antiseptic solution
- 18- to 20-gauge angiocatheter
- Intravenous extension tubing
- Three-way stopcock
- 10- and 20-mL syringes

B. Technique

- Prepare skin of appropriate hemithorax with antiseptic.
- Connect male end of three-way stopcock to female end of intravenous extension tubing. Connect syringe to three-way stopcock.
- Insert angiocatheter at point that is
 - at a 45-degree angle to skin, directed cephalad
 - $\circ~$ in second, fourth, or fifth intercostal space, just over top of rib, well above or below the areolar of the breast
 - in midclavicular line (Fig. 36.23A)
- As angiocatheter enters pleural space, decrease angle to 15 degrees above chest wall and slide cannula in while removing stylet (Fig. 36.23A).
- Attach male end of intravenous extension tubing to angiocatheter, open stopcock, and evacuate air with syringe (Fig. 36.23B).
- Continue evacuation as patient's condition warrants while preparing for permanent tube placement.
- Cover insertion site with petroleum gauze and small dressing after procedure.

Diagnostic Tap of Pleural Fluid

Follow the procedure for the insertion of a posterior chest tube, with the following differences.

A. Differences

- Use an angiocatheter, 20 gauge.
- Position patient without elevating side of fluid collection. It will be necessary to turn the affected side down only if quantity of fluid is small.
- Select insertion site in anterior or midaxillary lines below breast tissue for diffuse pleural collections. Direct catheter posteriorly after penetrating into pleural space.
- Keep system closed to prevent leakage of air into pleural space.

Anterior Mediastinal Drainage

Most mediastinal air collections cause only mild symptoms and are not under enough tension to require drainage. Their presence often precedes tension pneumothorax in the presence of lung disease and positive-pressure ventilation. Posterior mediastinal tube insertion, as described in the literature (42), is rarely required.

A. Indications

- Significant air accumulation with physiologic compromise (43)
 - Increased intracranial pressure (44)
 - Poor cardiac output because of impeded venous return
 - Critical interference with artificial ventilation
 - Competition with lungs for thoracic volume
 - Negative effect on pulmonary compliance
- Drainage of fluid
 - Mediastinitis after esophageal perforation
 - Postoperative

B. Contraindications

None absolute

C. Equipment

- Transillumination device with sterile transparent bag to cover tip
- Antiseptic for skin preparation
- Sterile gauze pads
- Sterile aperture drapes
- Surgical gloves
- 11 surgical blade
- Local anesthetic, as required
- Curved mosquito hemostat
- Drainage tube (see equipment for emergency evacuation of air leaks)
 - 10-Fr soft thoracostomy tube
 - Intravenous cannula system
 - 14- to 16-gauge angiocatheter
 - Intravenous extension tubing
 - Three-way stopcock
- 10- to 20-mL syringe
- $\hat{A}^{1/2}$ -in adhesive tape
- 4-0 nonabsorbable suture on small cutting needle with needle holder
- Connecting tubing and underwater suction device for indwelling tube

D. Precautions and Complications

The problems encountered in evacuating material from the mediastinum are similar to those encountered in placement of chest tubes. In contrast to tension pneumothorax, mediastinal collections tend to accumulate more gradually. For this reason, careful preparation of the patient and use of sterile technique are possible and essential. Refer to E and G under Thoracostomy Tubes at the beginning of this chapter for precautions and complications.

E. Technique

Drainage for longer than 12 hours normally dictates placing a 10- to 12-Fr tube by direct dissection because smaller tubes occlude readily. Select indwelling tubes only in the presence of significant lung disease or mediastinitis, where continued accumulations are anticipated. Remove the tubes as soon as possible because of the risks for infection.

Soft Mediastinal Tube Insertion

- Follow sterile technique throughout.
- Monitor infant for vital signs and oxygenation.



FIG. 36.24. Sequential radiographs. A: Tension pneumomediastinum (arrows). A mediastinal collection this massive is unusual. B: Successful drainage tube (arrow). C: The apparent slipping of the mediastinal cannula (arrow) is an artifact of patient rotation on this lateral view. There is still mediastinal air superiorly, but there was no patient compromise at this time.

- Determine by transillumination or radiograph the region of maximal mediastinal air accumulation (Fig. 36.24).
- Cover tip of transillumination light with sterile, clear plastic bag for use after skin preparation.
- Cleanse skin with antiseptic.
- Drape patient with aperture drape without obscuring infant.
- Infiltrate insertion site with 0.25 mL of local anesthetic.
- With no. 11 blade, make small stab wound through skin at subxiphoid.
- Using curved mosquito hemostat, dissect in the midline at 30-degree angle to chest wall in cephalad direction until entering mediastinal space. The mediastinum under tension should bulge downward.
- Insert soft chest tube into dissected tunnel, and direct tube cephalad and toward area of maximal transillumination.
- Observe tube for air rush or condensation while completing insertion. If loculations are evident, break them up with blunt dissection.
- Connect to closed drainage system at vacuum of 5 cm H_2O , and increase to 10 cm H_2O if necessary.

Accumulation in mediastinum is usually relatively slow; therefore, lower suction pressures are effective.

- Use low pressure to keep tube side holes patent while clearing air collection.
- Monitor efficacy by radiograph and transillumination (Fig. 36.24).
- Secure tube with suture, and tape as for thoracostomy tubes.

- If drainage stops with significant accumulation still evident on transillumination or radiograph:
 - Verify that accumulation is in mediastinum by lateral decubitus and lateral radiographs.
 - Verify tube position on radiographs.
 - Rotate tube.
 - Aspirate, but do not irrigate, tube; reattach to continuous drainage.
 - Change position of infant to move air toward tube.

Temporary Mediastinal Drainage with Intravenous Cannula

- Assemble equipment and prepare patient as for insertion by mediastinal dissection.
- Make a small stab wound in subxiphoid notch.

Mediastinal air under tension should be located in this area, pushing the liver and heart away from needle tip.

- Insert cannula with stylet at 45-degree angle to chest wall in cephalad direction.
- As soon as cannula passes through skin, lower cannula to 30 degrees from skin.
- Remove stylet, and attach connecting tubing, stopcock, and syringe.
- Advance cannula into mediastinal space cephalad and medially but toward area of maximal transillumination. Aspirate while advancing, and monitor cardiac tracing. Stop insertion if there is resistance, blood, or arrhythmia.
- Secure cannula in effective position, and attach intravenous extension tubing to underwater drainage system with suction pressure of 10 cm H₂O. The smaller cannula will require higher suction pressures unless the air accumulates slowly. Because air loculates within the mediastinum and the side holes occlude easily, small catheters are rarely effective for anything other than acute relief of tension. Remove cannula as soon as possible.

References

1. Jung AL, Nelson J, Jenkins MB, et al. Clinical evaluation of a new chest tube used in neonates. Clin Pediatr. 1991;30:85.

2. Lawless S, Orr R, Killian A, et al. New pigtail catheter for pleural drainage in pediatric patients. Crit Care Med. 1989;17:173.

3. Wood B, Dubik M. A new device for pleural drainage in newborn infants. Pediatrics. 1995;96:955.

4. Rothberg AD, Marks KH, Maisels MJ. Understanding the Pleurevac. Pediatrics. 1981;67:482.

5. Gonzalez F, Harris T, Black P, et al. Decreased gas flow through pneumothoraces in neonates receiving high-frequency jet versus conventional ventilation. J Pediatr. 1987; 110:464.

6. MacDonald MG. Thoracostomy in the neonate: a blunt discussion. NeoReviews. 2004;5:c301.

7. Moscowitz PS, Griscom NT. The medial pneumothorax. Radiology. 1976;120:143.

8. Dennis J, Eigen H, Ballantine T, et al. The relationship between peak inspiratory pressure and positive end expiratory pressure on the volume of air lost through a bronchopleural fistula. J Pediatr Surg. 1980;15:971.

9. Zidulka A, Braidy TF, Rissi MC, et al. Position may stop pneumothorax progression in dogs. Am Rev Respir Dis. 1982; 126:51.

10. Primhak RA. Factors associated with pulmonary air leak in premature infants receiving mechanical ventilation. J Pediatr. 1983;102:764.

11. Ryan CA, Barrington KJ, Phillips HJ, et al. Contralateral pneumothoraces in the newborn: incidence and predisposing factors. Pediatrics. 1987;79:417.

12. Kuhns LR, Bednarek FJ, Wyman ML. Diagnosis of pneumothorax or pneumomediastinum in the neonate by transillumination. Pediatrics. 1975;56:355.

13. Wyman ML, Kuhns LR. Accuracy of transillumination in the recognition of pneumothorax and pneumomediastinum in the neonate. Clin Pediatr. 1977;16:323.

14. Albelda SM, Gefter WB, Kelley MA, et al. Ventilator-induced subpleural air cysts: clinical, radiographic, and pathologic significance. Am Rev Respir Dis. 1983;127:360.

15. Allen RW, Jung AL, Lester PD. Effectiveness of chest tube evacuation of pneumothorax in neonates. J Pediatr. 1981;99: 629.

16. Batton DG, Hellmann J, Nardis EE. Effect of pneumothorax-induced systemic blood pressure alterations on the cerebral circulation in newborn dogs. Pediatrics. 1984;74:350.

17. Cartlidge PHT, Fox PE, Rutter N. The scars of newborn intensive care. Early Hum Dev. 1990;21:1.

18. Bhatia J, Mathew OP. Resolution of pneumothorax in neonates. Crit Care Med. 1985;13:417.

19. Merenstein GB, Dougherty K, Lewis A. Early detection of pneumothorax by oscilloscope monitor in the newborn infant. J Pediatr. 1972;80:98.

20. Noack G, Freyschuss V. The early detection of pneumothorax with transthoracic impedance in newborn infants. Acta Paediatr Scand. 1977;66:677.

Swischuk LE. Two lesser known but useful signs of neonatal pneumothorax. AJR. 1976;127:623.
 Genc A, Ozcan C, Erdener A, et al. Management of pneumothorax in children. J Cardiovasc Surg.

1998;39:849.

23. Mehrabani D, Kopelman AE. Chest tube insertion: a simplified technique. Pediatrics. 1989;83:784.

24. Mauer JR, Friedman PJ, Wing VW. Thoracostomy tube in an interlobar fissure: radiologic recognition of a potential problem. AJR. 1981;139:1155.

25. Strife JL, Smith P, Dunbar JS, et al. Chest tube perforation of the lung in premature infants: radiographic recognition. AJR. 1983;141:73.

26. Bowen A, Zarabi M. Radiographic clues to chest tube perforation of neonatal lung. Am J Perinatol. 1985;2:43.

27. McArtor RD, Saunders BS. Iatrogenic second-degree burn caused by a transilluminator. Pediatrics. 1979;63:422.

28. Moessinger AC, Driscoll JM, Wigger HJ. High incidence of lung perforation by chest tube in neonatal pneumothorax. J Pediatr. 1978;92:635.

29. Banagle RC, Outerbridge EW, Aranda JV. Lung perforation: a complication of chest tube insertion in neonatal pneumothorax. J Pediatr. 1979;94:973.

30. Jung A, Minton S, Roan Y. Pulmonary hemorrhage secondary to chest tube placement for pneumothorax in neonates. Clin Pediatr (Phila). 1980;19:624.

31. Kumar SP, Belik J. Chylothorax—a complication of chest tube placement in a neonate. Crit Care Med. 1984;12:411.

32. Rosegger H, Fritsch G. Horner's syndrome after treatment of tension pneumothorax with tube thoracostomy in a newborn infant. Eur J Pediatr. 1980;133:67.

33. Odita JC, Khan AS, Dincsoy M, et al. Neonatal phrenic nerve paralysis resulting from intercostal drainage of pneumothorax. Pediatr Radiol. 1992;22:379.

34. Arya H, Williams J, Ponsford SN, et al. Neonatal diaphragmatic paralysis caused by chest drains. Arch Dis Child. 1991;66:441.

35. Nahum E, Ben-Ari J, Schonfeld T, Horev G. Acute diaphragmatic paralysis caused by chest-tube trauma to phrenic nerve. Pediatr Radiol. 2001;31:444.

36. Yeh TF, Pildes RS, Salem MR. Treatment of persistent tension pneumothorax in a neonate by selective bronchial intubation. Anesthesiology. 1978;49:37.

37. Grosfeld JL, Lemons JL, Ballantine TVN, et al. Emergency thoracostomy for acquired bronchopleural fistula in the premature infant with respiratory distress. J Pediatr Surg. 1980; 15:416.

38. Faix RG, Naglie RA, Barr M. Intrapleural inoculation of Candida in an infant with congenital cutaneous candidiasis. Am J Perinatol. 1986;3:119.

39. Gooding C, Kerlan R Jr, Brasch R. Partial aortic obstruction produced by a thoracostomy tube. J Pediatr. 1981;98:471.

40. Ragosta KG, Fuhrman BP, Howland DF. Flow characteristics of thoracotomy tubes used in infants. Crit Care Med. 1990; 18:662.

41. Arda IS, Gurakan B, Aliefendioglu D, Tuzun M. Treatment of pneumothorax in newborns: use of venous catheter versus chest tube. Pediatr Int. 2002;44:78.

42. Purohit DM, Lorenzo RL, Smith CE, et al. Bronchial laceration in a newborn with persistent posterior pneumomediastinum. J Pediatr Surg. 1985;20:82.

43. Moore JT, Wayne ER, Hanson J. Malignant pneumomediastinum: successful tube mediastinostomy in the neonate. Am J Surg. 1987;154:687.

44. Tyler DC, Redding G, Hall D, et al. Increased intracranial pressure: an indication to decompress a tension pneumomediastinum. Crit Care Med. 1984;12:467.

37-Pericardiocentesis

Alan Benheim

A. Definitions

- Pericardium
 - A double layer of mesothelial lining surrounding the heart, consisting of the visceral pericardium on the epicardial surface and the parietal pericardium as an outer layer
 - Between the two layers there is normally a small amount of pericardial fluid (typically <5 mL for a neonate) that is thought to reduce friction.
- Pneumopericardium
 - Collection of air in the pericardial space
- Pericardial effusion
 - Accumulation of excess fluid in the pericardial space
- Pericardiocentesis
 - A procedure to remove air or excess fluid from the pericardial space, usually through a needle, small cannula, or drainage catheter
- Pericardial drain
 - A catheter or other drainage device left in place to allow intermittent or continuous evacuation of air or fluid from the pericardial space
 - Placed in select situations with recurring accumulation of air or fluid in the pericardial space
- Tamponade
 - Clinical condition with limited cardiac output because of external restriction of expansion of the heart, preventing normal cardiac filling, resulting in a decreased stroke volume and impaired cardiac output
 - \circ May be caused by:
 - Fluid or air in the pericardial space
 - Abnormalities of the pericardium (restrictive or constrictive)
 - Increased intrathoracic pressure associated with obstructive airway lung disease or tension pneumothorax
- Pulsus paradoxus (Fig. 37.1)
 - Respiratory variation in blood pressure, with a decrease in systolic blood pressure during inspiration during spontaneous respiration. (During positive-pressure ventilation, this is reversed, with a rise in systolic pressure during inspiration.)
 - This finding occurs during tamponade.
- B. Purpose
 - To evacuate air to relieve cardiac tamponade
 - To evacuate fluid to relieve cardiac tamponade

• To obtain fluid for diagnostic studies

C. Background

- The heart lies within a closed space, covered by the pericardium, which consists of an inner visceral layer and an outer parietal layer. The pericardial space is between these two layers. Ordinarily, the pericardial space of a neonate has <5 mL of pericardial fluid. This fluid acts as a lubricant to allow the heart to move with minimal friction. If the pericardial space fills with excess fluid, or if air accumulates, then the heart itself has less space available, and the pressure within the pericardium increases. This restricts venous return and impairs cardiac filling. This decrease in venous return and cardiac filling results in a reduced cardiac output. This clinical situation is called cardiac tamponade (1, 2, 3, 4 and 5).
- Neonates are at risk for cardiac tamponade
 - Accumulation of air dissecting into the pericardium from the respiratory system (Fig. 37.2) (4, 5, 6 and 7)
 - Pericardial fluid accumulation due to perforation or transudate from umbilical or percutaneous central venous catheter (Fig. 37.3) (1, 8, 9, 10, 11 and 12)
 - Cannulation for extracorporeal membrane oxygenation (13, 14)
 - Cardiac catheterization, either diagnostic or therapeutic (15)
 - Postoperative pericardial hemorrhage following cardiac surgery (2, 16)
 - Postpericardiotomy syndrome, typically 1 to 3 weeks after cardiac surgery (2, 16, 17)
 - Pericardial effusion as part of generalized edema/hydrops (3, 16)
 - Pericardial effusions due to infectious or autoimmune causes are less common in neonates than in older children.
- Clinical signs of cardiac tamponade may evolve gradually or rapidly (1, 3, 18).
- The primary therapy for cardiac tamponade is to evacuate the pericardial space; volume and pressors may be of transient benefit, but do not usually result in sustained clinical improvement (1, 8, 12, 16, 19).
- Cardiac tamponade may require urgent treatment with pericardiocentesis in infants with severe hemodynamic compromise (1, 16, 17).



FIG. 37.1. Pulsus paradoxus.

D. Indications (1, 12, 15, 16 and 17)

- Cardiac tamponade due to pneumopericardium
- Cardiac tamponade due to pericardial fluid
- Aspiration of pericardial fluid for diagnostic studies



FIG. 37.2. Chest radiograph with pneumopericardium.

E. Contraindications

- There are no absolute contraindications to performing pericardiocentesis in the setting of cardiac tamponade.
- Relative contraindication for diagnostic pericardiocentesis
 - Coagulopathy
 - Active infection. (However, infection may also be an indication for diagnostic pericardiocentesis in some clinical situations.)

F. Limitations

- Cannot readily evacuate thrombus
- Cannot remove mass lesions



FIG. 37.3. Echocardiogram image of preterm infant with pericardial effusion and central venous line in left atrium.

G. Equipment

- Antiseptic solution
- Sterile field with aperture drape or multiple drapes to be arranged around access site
- Sterile swabs or gauze pads
- Sterile gloves
- Local anesthetic, as needed
- 16- to 20-gauge intravenous cannula over 1- to 2-in needle
- Indwelling drainage catheter (optional)
- Three-way stopcock
- Short intravenous extension tubing (optional)
- 10- to 20-mL syringes
- Preassembled closed drainage system as for Emergency Evacuation of Air Leaks, Thoracostomy Tubes (optional)
- Connecting tubing and underwater seal for indwelling drain (optional)
- Transillumination device (optional, for pneumopericardium)
- Echocardiogram/sonography imaging device (optional in urgent situations)
- Specimen containers for laboratory studies, if procedure is diagnostic

H. Precautions

• Draining a large volume from the pericardial space can alter cardiac preloading conditions significantly, and some infants may require a supplemental intravascular fluid bolus after the pericardium is drained.

I. Techniques

- If ultrasound/echocardiographic imaging is available, and if time permits, then imaging can be performed to determine an optimal entry site and angle. In addition, the approximate distance required to reach the pericardial space can be estimated (15). Even after a sterile field is created, ultrasound imaging can be performed from a nonsterile area of the chest to monitor the effusion during the procedure. If imaging is done from a part of the sterile field, the transducer can be placed in a sterile sheath (or even a sterile glove), if needed. Care should be taken to avoid moving a probe with sterile cover back and forth between sterile and nonsterile areas.
- Similarly, evaluation with transillumination can be performed in cases of pneumopericardium, if time permits.
- Cleanse skin over xiphoid, precordium, and epigastric area with antiseptic. Allow to dry.
- Arrange sterile drapes, leaving the subxiphoid area exposed.
- Local anesthesia should be administered for the conscious patient. A typical example is 0.25 to 1.0 mL of subcutaneous 1% lidocaine instilled within 1 to 2 cm of the xiphoid process.
- Assemble the needle/cannula, three-way stopcock, and syringe so that the stopcock is open to both the needle and the syringe, but closed to the remaining port.
- The usual entry point in an infant is 0.5 to 1 cm below the tip of the xiphoid process, in the midline or slightly (0.5 cm) to the left of the midline. The needle should be elevated 30 to 40 degrees at the skin, and the tip should be directed toward the left shoulder (Fig. 37.4). A different approach may be used in certain cases, for example, if an echocardiogram suggests that most of the fluid is right-sided or apical.
- While advancing the needle, apply gentle negative pressure with the syringe. Continue advancing until air or fluid is obtained. If the syringe fills, use the third port of the stopcock to empty the syringe, or to attach a second syringe, and then aspirate more, repeating as needed. If diagnostic studies are desired, the fluid should be transferred to appropriate specimen containers.
 - If bloody fluid is aspirated, there could be a serosanguineous or hemorrhagic effusion, or the needle might have entered the heart (usually the right ventricle). There are a few clues that can be helpful in determining whether the needle has entered the heart; see J.



FIG. 37.4. Insertion of needle/cannula attached to three-way stopcock, in the subxiphoid space, directed toward the left shoulder.



FIG. 37.5. Echocardiogram images of pericardiocentesis. A: Echocardiogram image of pericardial effusion. B: Tip of needle in pericardial space. C: Pericardial effusion partially drained.

- A rhythmic tug, corresponding to the heart rate, may be felt as the needle enters the pericardium. Although this tugging sensation can reflect entering the myocardium, it can also be felt while the tip of the needle is positioned correctly within the pericardial space, and it does not necessarily mean that the needle has entered the heart.
- If ultrasound imaging is available, needle position can be determined either by visualizing the tip of the needle within the pericardial space or by demonstrating that the amount of pericardial fluid is diminishing as fluid is aspirated (Fig. 37.5). Some authors have described reinfusing a small amount of the aspirated fluid while imaging, to observe the location of microcavitation echoes (15, 20, 21).
- Once the needle is in the pericardial space, as much pericardial fluid or air should be evacuated as possible. To accomplish this, fix the needle in position and advance the cannula over the needle into the pericardial space. Remove the needle, and connect the cannula to a closed system for aspiration, such as a three-way stopcock and a syringe. Aspirate as much fluid/air as possible.
 - Note that small single-lumen catheters may easily become blocked.
 - A decision will need to be made whether to leave the cannula in place for any length of time or to remove it once the pericardium has been drained. This decision will vary in individual cases, but factors to consider include the likelihood of reaccumulation and need for repeat drainage versus the risk of infection or entry of free air with an indwelling cannula.
 - In certain cases, the operator may elect to evacuate the pericardial space directly through the needle, rather than placing a cannula.

J. Special Circumstances

- If ultrasound imaging is available, it may be helpful in planning the needle entry site and angle, as well as anticipating the distance required to reach the pericardial space (2, 15, 17, 20, 21).
- If transillumination is positive for free air before the procedure, it can be used to assess the adequacy of air evacuation after the procedure, and to look for evidence of reaccumulation. Because pneumothorax and pneumomediastinum are potential complications, the availability of transillumination may also be helpful after the procedure. Transillumination is not reliable to rule out free air or to distinguish between pericardial air and mediastinal air (5, 6).
- Upon initial aspiration of the pericardium, one may encounter several different things: air, serous fluid, serosanguineous or grossly bloody fluid, or fluid resembling infusate from a central line, including parenteral nutrients and lipid solutions. One should not be startled by any of these. Bloody fluid raises the concern that the needle may have entered the heart. Several clues may be helpful in distinguishing between pericardial fluid and intracardiac blood.
 - In an infant with tamponade, aspirating 10 mL of blood from the heart will have minimal effect on the acute hemodynamics, whereas draining as little as 5 to 15 mL from the pericardial space can result in significant hemodynamic improvement within 30 seconds.
 - If ultrasound is being used, the pericardial fluid volume will appear to be decreased if the needle is correctly positioned. In some cases, one can reliably identify the needle in the pericardial space (Fig. 37.5) (15).
 - Placing a few drops on a clean gauze may help distinguish the two sources, because serosanguineous fluid will separate into a central dark red zone and a more serous peripheral zone, but this can take several minutes.
 - Alternatively, a spun hematocrit can be performed rapidly if the unit has a readily available centrifuge; this also takes a few minutes.
- Draining a large volume from the pericardial space can alter cardiac preloading conditions significantly, and some infants may benefit from intravascular fluid boluses after the pericardium is drained.
- Pericardiocentesis is often an urgent or emergency procedure. The technique for pericardiocentesis described above applies when there is time for each step. In an infant with significant hemodynamic compromise, the operator may be forced to omit certain steps in the interest of time. This requires a judgment as to the baby's clinical status and the time delay involved for any given step, such as waiting for the ultrasound machine, preparing a larger sterile field, or assembling a three-way stopcock system. In extreme cases, this life-saving procedure might consist of pouring or swabbing Betadine over the subxiphoid area, followed by "blind†aspiration using any available needle and syringe, without anesthetic, and before any other equipment is available at the bedside (15).

K. Complications (15, 16 and 17,20,21)

- Pneumopericardium
- Pneumomediastinum
- Pneumothorax
- Cardiac perforation
- Arrhythmia
- Hypotension (if a large effusion is drained)

References

1. Nowlen TT, Rosenthal GL, Johnson GL, et al. Pericardial effusion and tamponade in infants with central catheters. Pediatrics. 2002;110:137.

2. Tsang TS, Barnes ME, Hayes SN, et al. Clinical and echocardiographic characteristics of significant pericardial effusions following cardiothoracic surgery and outcomes of echo-guided pericardiocentesis for management: Mayo clinic experience. 1979–1998. Chest. 1999;116:322.

3. Tamburro RF, Ring JC, Womback K. Detection of pulsus paradoxus associated with large pericardial effusions in pediatric patients by analysis of the pulse-oximetry waveform. Pediatrics. 2002;109:673.

4. Heckmann M, Lindner W, Pohlandt F. Tension pneumopericardium in a preterm infant without mechanical ventilation: a rare cause of cardiac arrest. Acta Paediatr. 1998;87:346.

5. Hook B, Hack M, Morrison S, et al. Pneumopericardium in very low birthweight infants. J Perinatol. 1995;15(1):27.

6. Cabatu EE, Brown EG. Thoracic transillumination: aid in the diagnosis and treatment of pneumopericardium. Pediatrics. 1979;64:958.

7. Bjorklund L, Lindroth M, Malmgren N, Warner A. Spontaneous pneumopericardium in an otherwise healthy full-term newborn. Acta Pediatr Scand. 1990;79:234.

8. Pesce C, Mercurella A, Musi L, et al. Fatal cardiac tamponade as a late complication of central venous catheterization: a case report. Eur J Pediatr Surg. 1999;9:113.

9. van Engelenburg KC, Festen C. Cardiac tamponade: a rare but life-threatening complication of central venous catheters in children. J Pediatr Surg. 1998;33:1822.

10. Fioravanti J, Buzzard CJ, Harris JP. Pericardial effusion and tamponade as a result of percutaneous silastic catheter use. Neonatal Network. 1998;17:39.

11. van Ditzhuyzen O, Ronayette D. Tamponnade cardiaque aprÃ"s catheterisme veineux central chez un nouveaune. Arch Pediatr. 1996;3:463.

12. Pezzati M, Filippi L, Chiti G, et al. Central venous catheters and cardiac tamponade in preterm infants. Intensive Care Med. 2004;30:2253.

13. Kurian MS, Reynolds ER, Humes RA, Klein MD. Cardiac tamponade caused by serous pericardial effusion in patients on extracorporeal membrane oxygenation. J Pediatr Surg. 1999;34:1311.

14. Becker JA, Short BL, Martin GR. Cardiovascular complications adversely affect survival during extracorporeal membrane oxygenation. Crit Care Med. 1998;26:1582.

15. Tsang TS, Freeman WK, Barnes ME, et al. Rescue echocardiographically guided pericardiocentesis for cardiac perforation complicating catheter-based procedures: The Mayo Clinic experience. J Am Coll Cardiol. 1998;32:1345.

16. Tsang TS, Oh JK, Seward JB. Diagnosis and management of cardiac tamponade in the era of echocardiography. Clin Cardiol. 1999;22:446.

17. Tsang TS, El-Najdawi EK, Seward JB, et al. Percutaneous echocardiographically guided pericardiocentesis in pediatric patients: evaluation of safety and efficacy. J Am Soc Echocardiogr. 1998;11:1072.

18. Berg RA. Pulsus paradoxus in the diagnosis and management of pneumopericardium in an infant. Crit Care Med. 1990;18:340.

19. Traen M, Schepens E, Laroche S, van Overmeire B. Cardiac tamponade and pericardial effusion due to venous umbilical catheterization. Acta Paediatr. 2005;94:626.

20. Muhler EG, Engelhardt W, von Bernuth G. Pericardial effusions in infants and children: injection of echo contrast medium enhances the safety of echocardiographically-guided pericardiocentesis. Cardiol Young. 1998;8:506.

21. Watzinger N, Brussee H, Fruhwald FM, et al. Pericardiocentesis guided by contrast echocardiography. Echocardiography. 1998;15:635.

38-Gastric and Transpyloric Tubes

Allison M. Greenleaf

A. Definitions

- Enteral feeding is defined as providing nutrients distal to the oral cavity (1).
- gastric tube is a tube inserted via the nose or mouth to the stomach (2).
- transpyloric tube is a tube passed through the pylorus to the duodenum or jejunum (2).

B. Purpose (3)

- To provide a route for feeding
- To administer medications
- To sample gastric or intestinal contents
- To decompress and empty the stomach

C. Background

- Coordinated suck and swallow does not develop until after 32 weeks' gestation (2).
- In preterm infants, enteral feedings stimulate the secretion of hormones that influence development of the gastrointestinal (GI) tract.
- Early introduction of feedings is associated with improved bone growth, glucose tolerance, bile flow, and lower risk of nosocomial infection (4).
- Early enteral feeding leads to lower cost and decreased length of hospital stay.
- Incorrect placement of gastric and transpyloric tubes (incidence as high as 21% to 43%) can lead to substantial morbidity and mortality (5,6 and 7).

Oral or Nasal Gastric Tubes

A. Indications (3,8)

- Poorly coordinated suck and swallow
- Abnormal gag reflex
- Insufficient oral intake
- Respiratory symptoms that prevent oral feeding

B. Contraindications

• Recent esophageal repair or perforation

C. Limitations

- Size of nares
- Type and amount of respiratory support
- Congenital anomalies of the nasopharynx

D. Equipment

- Suction equipment
- Cardiac monitor

- Infant feeding tube
 - \circ 3.5- or 5-French (Fr) feeding tube for infants weighing <1,000 g
 - \circ 5- or 8-Fr infant feeding tube for infants weighing >1,000 g
 - ¹/₂-in adhesive tape, pectin-based skin barrier
- Sterile water or saline
- 5- and 20-mL syringes
- Stethoscope
- Gloves
- pH paper

E. Precautions

- Measure and note appropriate length for insertion.
- Have suction apparatus readily available in case there is regurgitation.
- Do not push against any resistance. Perforation occurs even with very little force or sensation of resistance.
- Do not instill any material before verifying tube placement.
- Evaluate for possible esophageal perforation after insertion if there is:
 - Bloody aspirate
 - Increased oral secretion
 - Respiratory distress
 - \circ Pneumothorax
- Stop the procedure immediately if there is any respiratory compromise.
- Ensure that the open tube drains below the level of the infant's stomach.

TABLE 38.1 Guidelines for Minimum Orogastric Tube Insertion Length to Provide Adequate Intragastric Positioning in Very Low-Birthweight Infants

Weight (g)	Insertion Length (cm)
<750	13
750-999	15
1,000-1,249	16
1,250-1,500	17

Data from **Gallaher KJ**, **Cashwell S**, **Hall V**, et al. Orogastric tube insertion length in very low birth weight infants. *J Perinatol*. 1993;13:128.

F. Special Circumstances

- Feeding with umbilical catheters in situ should be done with caution, as there are insufficient data to guide practice (4,9).
- Tubing should be vented between feedings if continuous positive airway pressure (CPAP) is in place.
- The pH of gastric secretions does not appear to be affected by the method of feeding (continuous versus bolus). Administration of acid-reducing medications such as H_2 -receptor antagonists or proton-pump inhibitors does increase the pH of gastric secretions, but the pH is rarely >6 (6,10).

G. Technique

- Wash hands and put on gloves, maintaining aseptic technique.
- Clear infant's nose and oropharynx by gentle suctioning as necessary.
- Monitor infant's heart rate and observe for arrhythmia or respiratory distress throughout procedure.

- Position infant on back with head of bed elevated.
- Measure length for insertion by measuring distance from nose to ear to halfway between the xiphoid and umbilicus (4,6). Mark length on feeding tube with a loop of tape (Table 38.1) (11).
- Moisten end of tube with sterile water or saline.
- Oral insertion
 - Depress anterior portion of tongue with forefinger and stabilize head with free fingers.
 - Insert tube along finger to oropharynx.
- Nasal insertion (avoid this route in very low-birth-weight infants, in whom nasal tubes could cause periodic breathing and central apnea) (4).
 - Stabilize head. Elevate tip of nose to widen nostril.
 - Insert tip of tube, directing it toward occiput rather than toward vertex (Fig. 38.1).
 - Advance tube gently to oropharynx.
 - Observe for bradycardia.
- If possible, use pacifier to encourage sucking and swallowing.
- Tilt head forward slightly.
- Advance tube to predetermined depth.
 - Do not push against any resistance.
 - Stop procedure if there is onset of any respiratory distress, cough, struggling, apnea, bradycardia, or cyanosis.
- Determine location of tip. Injecting air to verify placement is not a reliable method, as the sound of air into the respiratory tract can be transmitted to the GI tract (4,6,12,13).
 - Aspirate any contents; describe and measure; determine acidity by pH paper.
 - Gastric contents may be clear, milky, tan, pale green, or blood-stained.
 - Gastric fluid pH should be <6 (5,6,10).
 - If the pH of the gastric aspirate is â%/¥6 or no aspirate is obtained, placement should be verified by radiography (5,14).
 - Suspect perforation or misplacement if no air or fluid is returned, or if there is onset of respiratory distress, blood in the tube, or difficult insertion.
- Secure indwelling tube to face with Â¹/₂-in tape. In preterm infants, apply the tape over a pectinbased barrier to prevent skin breakdown (15).
 - For feedings, attach to syringe.
 - For gravity drainage, attach specimen trap and position below level of stomach.
 - For decompression, a dual-lumen Replogle tube, connected to low continuous suction, is preferred.
- Pinch or close gastric tube during removal, to prevent emptying contents into pharynx.
- Document patient response, with any physiologic changes observed, and that tube placement was verified.



FIG. 38.1. Anatomic view of the neonatal nasopharynx. The natural direction in tube insertion is toward the nasal turbinates, where it might stop and give an impression of obstruction. By pushing the nostril up, one can direct a tube toward the occiput with less trauma.

H. Complications

- Apnea, bradycardia, or desaturation
- Obstruction of obligatory nasal airway
- Irritation and necrosis of nasal mucosa
 - Epistaxis
 - Ulceration
 - Perforation (Figs. 38.2 and 38.3) (16,17 and 18)
 - Posterior pharynx, particularly at level of cricopharnyx
 - Esophagus
 - Submucosal, remaining within mediastinum
 - Complete into thorax
 - Stomach
 - o Duodenum
- Misplacement on insertion (19)
 - Coiled in oropharynx (Fig. 38.4A)
 - Trachea (Fig. 38.4B)
 - Esophagus (Fig. 38.4C–E)
 - Eustachian tube (20)
- Displacement after insertion because of inappropriate length or fixation
 - Pulling back into esophagus
 - Prolapsing into duodenum
- Coiling and clogging of tube
- Grooved palate with long-term use of indwelling tube (4)
- Increased gastroesophageal reflux
- Infection (4,21,22)

Transpyloric Feeding Tube

A. Indications

- Severe gastroesophageal reflux with risk of aspiration
- Gastric distention with continuous positive airway pressure
- Delayed gastric emptying

- Gastric motility disorders
- Sampling of duodenojejunal contents
- Intolerance to gastric feeds

B. Contraindications

• Clinical condition that compromises duodenojejunal integrity: necrotizing enterocolitis, fulminant sepsis, shock, patent ductus arteriosus, recent small-bowel surgery



FIG. 38.2. Two radiographic views demonstrating a typical nasopharyngeal perforation with extrapleural malposition of a nasogastric tube at the right lung base (arrows). Either a traumatic endobronchial intubation or primary trauma from the nasogastric tube accounts for the perforation.

C. Limitations

• Long-term use may be associated with fat malabsorption, although recent studies suggest that there is no significant difference in growth over time (2,8).

D. Equipment

- Feeding tube
 - 3.5- or 5-Fr, 36-in silicone or Silastic tubes
 - Silastic tubes are preferred. Polyvinyl chloride tubes are not recommended for long-term use because the plasticizers are leached, stiffening the tube (4). "Nonstiffeningâ€ polyurethane tubes lose some of their compliance when left in place but stay softer for longer periods than polyvinyl chloride tubes.
- 20-mL syringes
- ¹/₂-in tape, pectin-based skin barrier
- pH paper
- Continuous-infusion pump and connecting tubing

E. Precautions

• When determining oral or nasal placement, individual assessment must be done to weigh the risks of compromising the nasal airway. Oral tubes are not at a significantly greater risk for dislodgement (23,24).
- Avoid pushing against any obstruction or resistance.
- Replace tubes as per manufacturer's recommendations. If the tube is stiff on removal, replace next tube sooner.
- If a tube has become partially dislodged, replace it rather than pushing it in farther.
- When using feedings that tend to coagulate in tubing, it may be necessary to flush the tube periodically with air or water.
- Use reliable infusion pumps that are safe for oral use by controlling rate and detecting obstructions.
- Limit infusion of hypertonic solutions and do not deliver bolus feedings beyond the pylorus.
- Consider the effect of continuous feedings on medication absorption.

F. Special Circumstances

• See oral or nasal gastric tubes, E.

G. Technique

- Insert orogastric tube as per gastric tubes above. Aspirate gastric contents.
- Measure distance from glabella to heels or from the nose to the ear to the xiphoid to the right lateral costal margin (14,25,26). Mark point with tape on transpyloric tube.



FIG. 38.3. Posterior perforation of the esophagus demonstrated in postmortem examination of 26week gestational-age infant. Upper probe is through perforation. Barium from a premortem study

spilled through the perforation, causing pleuritis. Perforation may have occurred as a result of endotracheal intubation, suctioning, or passage of gastric tube and is more common in smaller premature infants.

- Turn patient onto right side and elevate the head of the bed 30 to 45 degrees (6).
- Inject 10 mL/kg of air through orogastric tube to distend stomach. Close orogastric tube (14,27).
- Pass transpyloric tube to predetermined depth.
- After approximately 10 minutes with infant remaining on right side, gently aspirate through transpyloric tube. Tube may be in position within duodenum if aspirate is
 - Without air
 - Alkaline
 - Bilious (gold or yellow in color)
- Verify placement by checking pH of aspirate. If pH is >6 and the color of the aspirate is gold or yellow, the tube is likely in place and radiographic confirmation is not necessary. If no aspirate is obtained or the pH is $\hat{a}\%$, placement should be verified via radiography (5,10,14,28,29). The tip of the tube should be just beyond the second portion of duodenum (Fig. 38.5).
- Data are emerging regarding the use of bedside bilirubin testing as an added verification of placement. Studies suggest that intestinal bilirubin a% 45 mg/dL confirms placement (6,10,28).
- Avoid pushing to advance tube after initial placement. If tube is not in far enough, retape to give external slack and to allow peristalsis to carry tip to new position.
- Most often, if the tube does not cross the pylorus within the first $\hat{A}^{1/2}$ hour after passage, it is unlikely to pass in the next few hours, and it may be better to restart the procedure.
- After correct positioning, close transpyloric tube or start continuous infusion. Open gastric tube with syringe-barrel chimney or specimen trap to decompress stomach and to detect any transpyloric regurgitation.
- In larger infants, transpyloric tubes may be placed using a 6-Fr, unweighted, stylet-containing tube. This technique is not recommended for small infants.
- Transpyloric tubes may be placed with fluoroscopic guidance (6).

H. Complications (3) (See also Oral or Nasal Gastric Tubes, F)

- Misplacement into tracheobronchial tree, esophagus, or stomach, or incorrect position within proximal duodenum (19) (see Fig. 38.4).
 - \circ Aspiration
 - Pneumonia

The risk of aspiration with transpyloric feeding does not appear to be different from the risk with gastric feeding (8,24,30).

- Kinking or knotting of tube
- Hardening of polyvinyl chloride tube with leaching of bioavailable plasticizers
- Perforation of esophagus, stomach, duodenum (31,32)
- Perforation of kidney (33)
- Sepsis (21)
- Local infection (nasal colonization with staphylococci)
- Enterocolitis
 - Staphylococcus (21)
 - Necrotizing enterocolitis



FIG. 38.4. Radiographic examples of misplaced feeding tubes. A: Tube coiled in the oropharynx and upper esophagus, simulating an esophageal atresia. B: Tube into the left mainstem bronchus. C: Tube coiled in the lower esophagus. D: Tube doubled on itself in the stomach with its distal end in the esophagus (arrow). E: Tube only into the esophagus. A rush may be heard on auscultation over the stomach when air is injected through a tube lying in this position, making that an unreliable sign of gastric location.



FIG. 38.5. Radiographic demonstration of a transpyloric feeding tube that has passed the ligament of Treitz, well below the more appropriate level, increasing the risk of perforation or nutritional dumping.

- Development of pyloric stenosis (34)
- Formation of enterocutaneous fistula (35)
- Interference with absorption of medications (36)
- Malabsorption and GI disturbance (2,4,5,8)
 - Fat malabsorption with nasojejunal feeds
 - Dumping syndrome if hypertonic medications or feedings instilled too rapidly
 - Gastrointestinal disturbance as characterized by abdominal distention, gastric bleeding, and bilious vomiting
- Intussusception (37)

References

1. Directors ABO. Definitions of terms used in ASPEN guidelines and standards. J Parent Enter Nutr. 1995;19:1–2.

2. Groh-Wargo S, Thompson M, Cox JH, eds. Nutritional Care for High-Risk Newborns. 3rd ed. Chicago: Precept Press; 2000.

3. MacDonald PD, Skeoch CH, Carse H, et al. Randomized trial of continuous nasogastric, bolus nasogastric, and transpyloric feeding in infants of birth weight under 1400 g. Arch Dis Child. 1992;67:429.
4. Premji SS. Enteral feeding for high-risk neonates: a digest for nurses into putative risk and benefits to ensure safe and comfortable care. J Perinat Neonat Nurs. 2005;19:59–71.

5. Westhus N. Methods to test feeding tube placement in children. MCN Am J Matern Child Nurs. 2004;29:282–291.

6. Ellett MLC, Croffie JMB, Cohen MD, Perkins SM. Gastric tube placement in young children. Clin Nurs Res. 2005; 14:238–252.

7. Ellett ML. What is known about methods of correctly placing gastric tubes in adults and children. Gastroenterol Nurs. 2004;27:253–259.

8. McGuire W, McEwan P. Systematic review of transpyloric versus gastric tube feeding for preterm infants. Arch Dis Child Fetal Neonatol Ed. 2004;89:F245–F248.

9. Tiffany KF, Burke BL, Collins-Odoms C, Oelberg DG. Current practice regarding the enteral feeding of high-risk newborns with umbilical catheters in situ. Pediatrics. 2003; 112:20–23.

10. Metheny NA, Stewart BJ. Testing feeding tube placement during continuous tube feedings. Appl Nurs Res. 2002;15: 254–258.

11. Gallaher KJ, Cashwell S, Hall V, et al. Orogastric tube insertion length in very low birth weight infants. J Perinatol. 1993;13:128.

12. Huffman S, Pieper P, Jarczyk KS, et al. Methods to confirm feeding tube placement: application of research in practice. Pediatr Nurs. 2004;30:10–13.

13. Nyqvist KH, Sorell A, Ewald U. Litmus tests for verification of feeding tube location in infants: evaluation of their clinical use. J Clin Nurs. 2005;14:486–495.

14. Ellett MLC. Important facts about intestinal feeding tube placement. Gastroenterol Nurs. 2006;29:112–124.

15. Dollison EJ, Beckstrand J. Adhesive tape vs pectin-based barrier use in preterm infants. Neonatal Network. 1995;14:37.

16. Sapin E, Gumpert L, Bonnard A, et al. Iatrogenic pharyngoesophageal perforation in premature infants. Eur J Pediatr Surg. 2000;10:83.

17. Agarwala S, Dave S, Gupta AK, et al. Duodeno-renal fistula due to a nasogastric tube in a neonate. Pediatr Surg Int. 1998;14:102.

18. Mattar MS, Al-Alfy AA, Dahniya MH, et al. Urinary bladder perforation: an unusual complication of neonatal nasogastric tube feeding. Pediatr Radiol. 1997;27:858.

19. McWey RE, Curry NS, Schabel SI, et al. Complications of nasoenteric feeding tubes. Am J Surg. 1988;155:253.

20. Wynne D, Borg H, Geddes N, Fredericks B. Nasogastric tube misplacement into Eustachian tube. Int J Pediatr Otorhinolaryngol. 2003;67:185–187.

21. Matlow A, Wray R, Goldman C, et al. Microbial contamination of enteral feed administration sets in a pediatric institution. Am J Infect Control. 2003;31:49–53.

22. Lee JS, Polin RA. Treatment and prevention of necrotizing enterocolitis. Semin Neonatol. 2003;8:449–459.

23. Hawes J, McEwan P, McGuire W. Nasal versus oral route for placing feeding tubes in preterm or low birth weight infants. Cochrane Database Syst Rev. 2004;3:CD003952.

24. Meert KL, Daphtary KM, Metheny NA. Gastric vs small-bowel feeding in critically ill children receiving mechanical ventilation: a randomized controlled trial. Chest. 2004;126:872–878. P.298

25. Harrison AM, Clay B, Grant MJ, et al. Nonradiographic assessment of enteral feeding tube position. Crit Care Med. 1997;25:2055.

26. Chellis MF, Sanders SV, Dean JM, et al. Bedside transpyloric tube placement in the pediatric intensive care unit. J Parent Enter Nutr. 1996;20:88.

27. DaSilva PS, Paulo CS, de Oliveira Iglesias SB, et al. Bedside transpyloric tube placement in the pediatric intensive care unit: a modified insufflation air technique. Intensive Care Med. 2002;28:943–946.

28. Gharpure V, Meert KD, Sarnaik AP, et al. Indicators of postpyloric feeding tube placement in children. Crit Care Med. 2000;28:2962.

29. Metheny NA, Schneiker R, McGinnis J, et al. Indicators of tubesite during feedings. J Neurosci Nurs. 2005;37:320–325.

30. Lyons KA, Brilli RJ, Wieman RA, Jacobs BR. Continuation of transpyloric feeding during weaning of mechanical ventilation and tracheal extubation in children: a randomized controlled trial. J Parent Enteral Nutr. 2002;26:209–213.

31. Chen JW, Wong BWK. Intestinal complications of nasojejunal feeding in low birth-weight infants. J Pediatr. 1974; 85:109.

32. Flores JC, Lopez-Herce J, Sola I, Carrillo A. Duodenal perforation caused by a transpyloric tube in a critically ill infant. Nutrition. 2006;22:209–212.

33. Perez-Rodrigues J, Quero J, Frias EG, et al. Duodenal perforation by silicone rubber tube. J Pediatr. 1978;92:113.

34. Latchaw LA, Jacir NN, Harris BH. The development of pyloric stenosis during transpyloric feedings. J Pediatr Surg. 1989; 24:823.

35. Patrick CH, Goodin J, Fogarty J. Complication of prolonged transpyloric feeding: formation of an enterocutaneous fistula. J Pediatr Surg. 1988;23:1023.

36. Sneed RC, Morgan WT. Interference of oral phenytoin absorption by enteral tube feedings. Arch Phys Med Rehabil. 1988;69:682.

37. Hughes U, Connolly B. Small-bowel intussusceptions occurring around nasojejunal enteral tubesâ€"three cases occurring in children. Pediatr Radiol. 2001;31:456–457.

39-Gastrostomy

Thomas T. Sato

A. Indications

- Inability to swallow
 - Neurologic or neuromuscular deficit leading to uncoordinated oropharyngeal swallowing
 - Complex congenital malformations, including esophageal atresia or Pierre Robin sequence, not undergoing early correction
 - Administration of supplemental feedings despite normal swallowing but inadequate oral intake
 - Chronic disease states
 - Requirement for unpalatable diet or need for consistent glucose source (i.e., glycogen storage disease)
- Presence of anatomic intestinal anomalies or functional intestinal dysmotility disorders that prevent normal enteral feeding
- Gastric decompression, particularly when respiratory compromise makes prolonged presence of nasogastric/ orogastric tube undesirable
 - Chronic gastrointestinal decompression: duodenal atresia, gastric volvulus, gastric gas/bloat from antireflux procedure

• Emergency gastric decompression for inadequate mechanical ventilation secondary to gastric distention: tracheoesophageal fistula/esophageal atresia

B. Contraindications

The presence of treatable conditions that unduly increase operative risks, e.g., active infection or coagulopathy. These conditions should be treated aggressively prior to elective gastrostomy placement. Predictably, newborn infants with pure esophageal atresia or tracheoesophageal fistula with esophageal atresia have small stomach volumes (microgastria), making gastrostomy placement more difficult. C. Equipment

Operative Insertion

- Sterile neonatal laparotomy instruments
- 12- to 14- French (Fr) mushroom (de Pezzar or Malecot) single-lumen catheters, 14-Fr Ross balloontype gastrostomy catheters, or similar size latex-free Foley-type silastic catheters
- 3-0 or 4-0 absorbable sutures on taper needles
- Local infiltrative anesthesia (0.25% bupivicaine at 1 mL/kg bodyweight)

Maintenance Care after Insertion

- Fixation device
 - Modified feeding nipple, or
 - Cotton gauze bolster dressing
- Stomadhesive (Squibb, Princeton, NJ, USA)
- 3-0 or 4-0 permanent suture on cutting needle

D. Technique for Insertion

Neonatal gastrostomy placement often requires general anesthesia and frequently occurs as an adjunct to other abdominal operations. The technique varies according to individual surgeon preference, but the Stamm procedure described in 1894 is one of the most frequently used in premature infants and neonates (1). The gastrostomy should lie in the center of a triangle formed by the left costal margin, umbilicus, and xiphoid (Fig. 39.1).

Some data suggest that gastrostomy placement along the lesser curvature of the stomach may reduce the incidence of new-onset, postoperative gastroesophageal reflux (2,3). However, the small size of the neonatal stomach may make placement along the greater curvature more practical.

- After insertion of a nasogastric tube, sterile preparation of the skin, and delivery of intravenous antibiotics (typically a first-generation cephalosporin), enter the peritoneal cavity through either a transverse incision (Fig. 39.1) or a supraumbilical midline incision. Identify the stomach and elevate the anterolateral greater curvature of the stomach through the wound.
- Place two concentric, seromuscular purse-string sutures on the anterior greater curvature of the stomach body. Avoid injury to the gastroepiploic vessels (Fig. 39.2).
 - Inner purse-string for hemostasis
 - o Outer purse-string for inversion of mucosa and fixing of stomach to abdominal wall



FIG. 39.1. Landmarks for gastrostomy. The primary horizontal incision is left supraumbilical. The gastrostomy tube will pass through the abdomen at a separate site in the center of a triangle formed by the xiphoid, umbilicus, and left costal margin.

• Make a stab wound through stomach wall (gastrotomy) in the center of purse-string sutures. With a stylet or grooved director inside the catheter, flatten the mushroom catheter head and introduce catheter through gastrotomy. Verify position of tube inside stomach.



FIG. 39.2. Site for concentric sutures for Stamm procedure. Entrance into stomach is on greater curvature midway between esophagus and pylorus.



 \mathcal{GU} . FIG. 39.3. After the tube is secured inside the stomach and passed through a stab wound in the

abdominal wall, the anterior wall of the stomach is sutured to the inner wall of the abdomen.

- Tie sutures sequentially. The inner purse-string suture secures the stomach around the catheter and provides hemostasis. The outer suture allows for inversion and water-tight seal of the stomach wall.
- At the identified exit site, make a stab wound through the abdominal wall (Fig. 39.1). A 3-mm dermal punch is often useful for this maneuver. Insert a curved hemostat through the abdominal wall exit site and expose the undersurface of the abdominal wall through the incision. Secure the stomach to the abdominal wall by placing three or four absorbable sutures through the gastric wall and the abdominal wall (Fig. 39.3).
- With a hemostat, pull the gastrostomy tube through the abdominal wall exit site until the stomach is snug against the abdominal wall. Tie the previously placed sutures while placing gentle traction on the gastrostomy tube (Fig. 39.3).
- Secure the gastrostomy tube to the skin with a suture to prevent inadvertent removal. Document the length of the gastrostomy tube outside the abdomen. Close the abdominal incision in routine fashion.

E. Fixation of Tube after Surgical Placement

Attention to gastrostomy tube fixation on the abdominal wall is critical to encourage proper gastrocutaneous tract formation and to prevent inadvertent removal or site irritation. The gastrostomy tube should be kept perpendicular to the abdominal wall to maintain the smallest possible orifice and to minimize leakage. A dressing that supports the gastrostomy tube in this position for 5 to 7 postoperative days is important. Once the dressing is removed and the wound healed, the gastrostomy tube may be secured to the abdominal wall with porous tape to avoid maceration of the skin.

Feedings may begin on the first postoperative day and increase as tolerated. Avoid clamping the gastrostomy tube with initial feedings. Use gravity suspension to prevent loss of gastric contents while allowing decompression of the stomach during drip feeding. Bolus gastrostomy feeding is generally started once the goal enteral feeding volume has been established using continuous drip. Fixation of Gastrostomy to Abdominal Wall

• Catheter bridge (Fig. 39.4)

- Either latex or nonlatex catheters may be used.
- Cut a large, firm catheter at its wider end to a length of 3 to 4 cm.
- Cut opposing holes at the midpoint of the catheter bridge just wide enough to admit but not constrict the gastrostomy tube.
- Pull the gastrostomy tube through holes to create a tight seal on the catheter without constricting the lumen of the tube.



FIG. 39.4. Latex bridge at gastrostomy exit stabilizes tube perpendicular to skin, keeping stoma narrow to avoid leakage. Rotating the bridge around the tube allows change in contact points with the skin. Note how the flared end of the mushroom catheter is pulled to keep the stomach apposed to the abdominal wall.

- Insert distal end of catheter into gastrostomy site as described above. If using a Foley catheter, inject 2 to 4 mL of saline into balloon.
- Push catheter bridge snug against skin of abdominal wall.

Pull Foley catheter balloon or mushroom catheter flange against the stomach and abdominal wall.

- Gauze Bolster Dressing
 - Fold gauze pad into bolster and use Steri-Strips or tape to affix bolster to abdominal wall next to the gastrostomy tube.
 - Loop the gastrostomy tube over the bolster, maintaining the gastrostomy tube perpendicular to the abdominal wall.
 - Secure gastrostomy tube to bolster and abdominal wall.
- Modified, soft, feeding nipple (4) (Fig. 39.5)
 - Because it is critical to keep the site as dry as possible, there must be modification of the nipple for good circulation of air.
 - Excise an elliptical window in flared base of nipple in circumferential direction 1 to 1.5 cm in length.
 - Make 1-cm crosscut in tip of nipple.
 - Slide nipple over gastrostomy tube until there is contact with abdominal wall.
 - Adjust tension on tube to pull gastrostomy balloon or flange against gastric wall.



FIG. 39.5. Modified feeding nipple. The elliptical hole at the base allows air circulation and regular cleaning of the skin as important factors in avoiding maceration of the site. (From Kappell DA, Leape LL. A method of gastrostomy fixation. J Pediatr Surg. 1975;10:523, with permission.)

- Tape or suture nipple flange to skin. Clean skin around gastrostomy with cotton applicators and keep area under nipple dry to avoid skin maceration.
- F. Maintenance Care of Gastrostomy
 - Maintain contact between stomach and abdominal wall.
 - Prevent gastric distention.
 - Keep gastrostomy balloon or flange pulled snugly against stomach wall.
 - Avoid pressure necrosis of abdominal wall (Fig. 39.4).
 - Avoid inadvertent dislodgement of gastrostomy.
 - Keep tube immobile at insertion site to maintain stoma as small as possible.
 - Use careful fixation to maintain perpendicular position.

Keep some slack in the tube when it is suspended.

- Prevent migration of tube through pylorus or esophagus by:
 - Proper fixation
 - Comparing length of external tube with postoperative length
 - Observing for signs of obstruction, for example,
 - Gastric distention, feeding intolerance
 - Increased drainage from oral gastric or gastrostomy tube
 - Bilious drainage
 - New-onset or increased gastroesophageal reflux

- Minimize leak from gastrostomy site in long-term gastrostomies.
 - Maintain adequate fit of tube in stoma.
 - Avoid local infection.
 - Treat leaking gastrostomy early.
 - Remove tube for 24 hours to allow partial closure.
 - Replace mushroom catheter with Foley catheter, and pull balloon (inflated with 2 to 5 mL H₂O) against abdominal wall to seal leak. Use catheter bridge or tape to secure tube.
 - Apply Stomahesive around catheter to decrease excoriation and to encourage epithelialization. Change Stomahesive every 3 to 4 days, as long as seal remains.
 - Maintain perpendicular position of gastrostomy tube.
 - Keep tube unclamped.
 - Maintain skin and stoma hygiene.
 - Cleanse daily with soap and water; "crusty†exudate may be removed with half-strength hydrogen peroxide.
 - Keep area dry. Change dressing after cleansing skin and whenever wet.
 - Treatment of granulation tissue at gastrostomy site
 - Silver nitrate: Apply daily for 3 to 5 days.
 - 0.5% Triamcinolone ointment: Apply three times a day for 5 to 7 days.
 - Cautery: May require local or general anesthesia
- G. Replacing Gastrostomy Tubes
 - During the first 2 to 4 weeks after open gastrostomy placement, reoperation for replacement may be necessary.
 - After formation of gastrocutaneous fistula, nonoperative placement is possible.
 - To avoid stoma closure, replace within 4 to 6 hours.
 - Use deflated Foley catheter for replacement prior to formation of well-epithelialized tract.
 - If the catheter does not pass easily, a flexible guidewire may be placed through the tract and confirmed in gastric position by fluoroscopy.
 - Lubricate catheter generously with water-soluble lubricant, and insert gently.
 - Inflate balloon with 2 to 4 mL of water, and pull against stomach wall.
 - Secure with fixation device.
 - Mark outside length of catheter to help detect internal or external migration of balloon.

Prior to feeding, confirm placement of gastrostomy with water-soluble contrast study if replacement is difficult or uncertain.

- For replacement, use mushroom catheter tube or balloon-type gastrostomy tube in wellestablished tract.
 - Carefully determine direction of tract.
 - Lubricate catheter.
 - Stretch tip of mushroom catheter with introducer.
 - Apply gentle pressure to insert catheter. Avoid force, which may lead to traumatic separation of stomach from abdominal wall.
- \circ Confirm intragastric position by one of the following methods.
 - For recent gastrostomy
 - Instill 15 to 30 mL of water-soluble contrast and obtain decubitus radiograph. Remove contrast.
 - For well-established gastrostomy, aspirate for gastric contents. If there is any doubt, obtain contrast study prior to initiating feeding.

H. Discontinuation of Gastrostomy Tube (5) General

- Remove tube and apply gauze dressing.
- Allow spontaneous closure, usually 4 to 7 days, or
- Approximate skin edge with skin closure tape.

Persistent Gastrocutaneous Fistula

- Cause: Granulation and epithelialization of gastrocutaneous tract
 - Remove tube.
 - Cauterize granulation tissue or epithelium within stoma with silver nitrate sticks.
 - Seal orifice with Stomahesive.
 - Approximate edges with surgical tape.
- Persistent gastrocutaneous fistula of 4 to 6 weeks: Surgical closure required. If skin is becoming macerated, replace gastrostomy and use protective skin ointment prior to surgical closure.

I. Percutaneous Endoscopic Gastrostomy

The use of percutaneous endoscopic technique has become well established for primary gastrostomy tube placement in infants and children. There are several techniques, performed either with or without endoscopy.

- Percutaneous endoscopic gastrostomy (PEG)
 - "Pull†technique (6)
 - "Push†technique (7)
 - "Poke†technique (8)
- Nonendoscopic (9)
- Currently available instrumentation and fiber-optic endoscopes allow for the safe performance of percutaneous gastrostomy in selected neonates, infants, and children. Placement typically requires general anesthesia for airway protection and control. The original method of PEG described by Gauderer in 1980 involves four basic elements:
 - Endoscopic gastric insufflation to approximate stomach to abdominal wall
 - Percutaneous placement of an introducer into the stomach under direct endoscopic visualization
 - Placement of a guidewire through the introducer and retrieval of the guidewire with an endoscopic snare; removal of the endoscope and the snared guidewire through the infant's mouth
 - Attachment of a gastrostomy tube to the guidewire and pulling it through the infant's oropharynx, esophagus, and into the stomach. The gastrostomy tube is modified with a tapered end where it is pulled through the abdominal wall and an intragastric mushroom-type flange to keep it from pulling completely out of the stomach (10).

To provide prophylaxis against wound infection, a single intravenous dose of a firstgeneration cephalosporin or equivalent is recommended prior to PEG placement.

J. Laparoscopic Gastrostomy Placement

Placement of gastrostomy tubes in infants using a laparoscopic approach has been described as safe, efficacious, and may have a lower complication rate than percutaneous endoscopic gastrostomy (11). This approach allows direct visualization of the stomach via a 3- or 5-mm laparoscope.

- The laparoscope may be inserted either through the umbilicus or via a left upper quadrant incision at the proposed gastrostomy tube site.
- CO_2 insufflation to 8 to 10 torr is generally sufficient to visualize the peritoneal cavity.
- The stomach is directly visualized, grasped, and traction sutures placed to elevate the stomach to the abdominal wall; the gastrostomy tube is inserted into the stomach and the stomach is sutured to the abdominal wall.

K. Emergency Percutaneous Gastric Decompression

We describe an emergency percutaneous technique for gastric decompression that may be used as a lifesaving measure when there is either respiratory failure or a high probability of gastric rupture in the presence of extreme gastric distention.

The primary indication for this type of procedure is for infants with massive abdominal distention causing respiratory failure that cannot be decompressed using an orogastric or nasogastric tube, for example, in infants with esophageal atresia and unrepaired tracheoesophageal fistula, requiring emergency transport to a definitive care facility:

- Prepare skin in upper left abdomen with Betadine.
- If possible, transilluminate abdomen to verify position of distended stomach away from liver.
- Make a small weal with 1% lidocaine to provide local anesthesia.
- Using a 20-gauge catheter with needle stylet, puncture the abdominal wall at the junction of the left anterior rib cage and the lateral border of the rectus abdominis muscle. Advance needle through wall into stomach.
- Remove needle and advance catheter into stomach. Attach short IV extension tubing, three-way stopcock, and syringe. Aspirate only enough air to relieve tamponade effect and improve ventilation. Avoid completely emptying stomach.
- Secure catheter and keep in place until surgical evaluation is possible.

L. Complications (10,12,13,14 and 15)

The complications associated with gastrostomy placement in the neonate may be characterized as either immediate or long-term (late). Most immediate complications are technical or mechanical in nature. Immediate Complications (in approximate order of frequency)

- Pneumoperitoneum (after PEG)
- Wound infection, dehiscence, ventral hernia
- Prolonged ileus, gastric atony, feeding intolerance
- Separation of stomach from anterior abdominal wall
- Intraperitoneal spillage, gastric leak, peritonitis
- Injury to posterior wall of stomach on initial insertion or reinsertion
- Perforation or injury to other organs
 - o Diaphragm
 - o Esophagus (16)
 - \circ Liver, spleen
 - $\circ \quad Colon$

Late Complications:

- Dislodgement
 - Inadvertent removal
 - Internal or external migration (17)
 - Intraperitoneal, extragastric placement

- Catheter deterioration
 - Catheter fracture
 - Balloon rupture
- Tube occlusion
- Persistent leak
 - Wound breakdown
 - Granulation tissue and skin irritation
 - Infection, particularly Candida
 - Loose gastrostomy with leakage
 - Electrolyte imbalance
 - \circ Malnutrition
- New-onset or worsening gastroesophageal reflux disease (18)
- Persistent gastrocutaneous fistula following removal
- Adhesive bowel obstruction
- Prolapse of gastric mucosa
 - Bleeding
 - Excessive leakage
- Gastric torsion around catheter

References

1. Stamm M. Gastrostomy by a new method. Med News (NY). 1894;65:324.

2. Stringel G. Gastrostomy with antireflux properties. J Pediatr Surg. 1990;25:1019.

3. Seekri IK, Rescoria FJ, Canal DF, et al. Lesser curvature gastrostomy reduces the incidence of postoperative gastroesophageal reflux. J Pediatr Surg. 1991;26:982.

4. Kappel DA, Leape LL. A method of gastrostomy fixation. J Pediatr Surg. 1975;10:523.

5. Ducharme JC, Youseff S, Tilkin F. Gastrostomy closure: a quick, easy and safe method. J Pediatr Surg. 1977;12:729.

6. Gauderer MWL, Ponsky JL, Izant RJ Jr. Gastrostomy without laparotomy: a percutaneous endoscopic technique. J Pediatr Surg. 1980;15:872.

7. Sachs BA, Vine HS, Palestrant AM, et al. A non-operative technique for establishment of a gastrostomy in the dog. Invest Radiol. 1983;18:485.

8. Russell TR, Brotman M, Norris F. Percutaneous gastrostomy. A new simplified and cost-effective technique. Am J Surg. 1984; 148:132.

9. Cory DA, Fitzgerald JF, Cohen MD. Percutaneous nonendoscopic gastrostomy in children. AJR. 1988;151:995.

10. Gauderer MWL, Stellato TA. Gastrostomies: evolution, techniques, indications and complications. Curr Probl Surg. 1986; 23:661.

11. Zamakhshary M, Jamal M, Blair GK, et al. Laparoscopic vs percutaneous endoscopic gastrostomy tube insertion: a new pediatric gold standard? J Pediatr Surg. 2005;40:859.

12. Gallagher MW, Tyson KRT, Aschcraft KW. Gastrostomy in pediatric patients: an analysis of complications and techniques. Surgery. 1973;536:74.

13. Cywes S. Stomas in children. S Afr Med J. 1976;50:815.

14. Campbell JR, Sasaki TM. Gastrostomy in infants and children: an analysis of complications and techniques. Am Surg. 1974; 40:505.

15. Gauderer MWL. Percutaneous endoscopic gastrostomy: a 10-year experience with 220 children. J Pediatr Surg. 1991;26:288.

16. Kenigsberg K, Levenbrown J. Esophageal perforation secondary to gastrostomy tube replacement. J Pediatr Surg. 1986;21:946.

17. Currarino G, Votteler T. Prolapse of the gastrostomy catheter in children. AJR, Radium Ther Nucl Med. 1975;123:737.

18. Jolley SG, Tunnel WB, Hoelzer DJ, et al. Lower esophageal pressure changes with tube gastrostomy: a causative factor of gastroesophageal reflux in children? J Pediatr Surg. 1986;21:624.

40-Neonatal Ostomy and Gastrostomy Care

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An ostomy is the construction of a permanent or temporary opening in the intestine (enterostomy) or urinary tract (urostomy) through the abdominal wall to provide fecal or urinary diversion, decompression, or evacuation (1). Gastrostomies (G tubes) are stomas that allow direct access into the stomach and are used for feeding, medication administration, and decompression. This chapter discusses care of simple and complex ileostomies, colostomies, urostomies, and gastrostomies. Tracheostomy care is discussed in Chapter 35.

Enterostomies and Urostomies in the Neonate

A. Indications

Ostomies may be indicated in the neonate for a variety of congenital or acquired conditions (Table 40.1). The stoma is usually temporary, and reanastomosis of the bowel or urinary tract and closure of the stoma is performed during infancy or early childhood (2,3).

B. Types of Ostomies

- There are several types of intestinal stomas. The patient's condition, the segment of bowel affected, and the size of the patient's abdomen often determine the type of stoma and its external location. Figure 40.1 depicts the most common types of neonatal stomas (1).
- Urostomies are urinary diversions constructed to bypass a dysfunctional portion of the urinary tract. Ileal conduits and ureterostomies are rarely performed in the neonatal period.
- A vesicostomy is an opening directly from the bladder through the abdominal wall and is a more common urinary diversion in the neonate. Urine flows freely through the stoma from the bladder.

C. Ostomy Assessment

The neonate with a stoma needs careful observation and assessment for a variety of potential complications (4). Monitoring the infant for function of the ostomy is paramount in the initial postoperative period. Possible surgical complications are paralytic ileus, intestinal obstruction, anastomic leak, and stomal necrosis. The factors to be considered during evaluation of the stoma are listed below.

- Type of stoma: The segment of bowel from which the stoma is made.
- Viability: A healthy stoma should be bright pink to beefy red and moist, indicating adequate perfusion and hydration (Fig. 40.2). The stoma is formed from the intestine, which is very vascular and therefore may bleed slightly when touched or manipulated, but the bleeding usually resolves quickly. The stoma is not sensitive to touch because it does not have somatic afferent nerve endings (4).
 - A purple or dark brown to black stoma with loss of tissue turgor and dryness of the mucous membrane may indicate ischemia and possible stomal necrosis.
 - A pale pink stoma is indicative of anemia.
- Size: The stoma shape (round, oval, mushroom, or irregular) and diameter (length and width) in inches or millimeters is noted. In the early postoperative period, the stoma will be edematous. After

the first 48 to 72 hours, the edema should resolve and result in a reduction in size of the stoma, which should, however, still remain everted from the skin surface. Stomas generally continue to decrease in size over 6 to 8 weeks postoperatively. It is not uncommon for the stoma to become edematous when exposed to air while changing the pouch; this edema generally resolves quickly when the pouch is replaced.

Stomal height: The degree of protrusion of stoma from the skin. Ideally, the surgeon will evert the • stoma prior to suturing it to the skin to produce an elevation, which will promote a better seal with the ostomy wafer. With the stoma elevated above the surface of the skin, the effluent will be more likely to go into the pouch instead of staying in contact with the skin (2). Eversion of the stoma, referred to as maturing the stoma, is not always possible in neonates, in whom blood supply may be tenuous, and in situations in which the bowel is markedly edematous (1,5).

TABLE 40.1 Conditions Necessitating Ostomy in the Neonate			
Disease/Congenital Anomaly	Most Common Location of Stoma		
Intestinal atresia	Duodenum, ileum, or jejunum		
Meconium ileus	Ileum		
Necrotizing enterocolitis	Ileum or jejunum		
Hirschsprung disease	Sigmoid colon		
Imperforate anus/anorectal malformations	sColon		
Volvulus	Ileum or jejunum		
Bladder extrophy	Bladder		

TADLE 40.1 Conditions Neoessitating Octo • 41 NT



FIG. 40.1. A: End stoma. The end of the bowel is everted at the skin surface. B: Loop stoma. Entire loop of bowel is brought to the skin surface and opened to create a proximal, or functioning, end and a distal, or nonfunctioning, end. The distal side is called a mucus fistula because of the normal mucus secretions it produces. C: Double-barrel stoma. Similar to a loop stoma, except the bowel is divided into two stomas, a proximal and a distal stoma. The distal stoma functions as a mucus fistula. (Adapted from Gauderer MWL. Stomas of the small and large intestine. In: O'Neil JA, Rowe MI, Grosfeld JL, et al., eds. Pediatric Surgery. 5th ed. St. Louis, MO: Mosby; 1998:1349–1359, with permission.)

- Stomal construction: The ostomy may be an end, loop, or double-barrel (Figs. 40.1 and 40.3).
- Abdominal location
- Peristomal skin: Ideally the peristomal skin should be intact, nonerythematous, and free from rashes. However, frequently the stoma(s) is not separate from the surgical incision (Fig. 40.4). There is often not enough space on the baby's abdomen for the surgeon to create separate incisions. In addition, stomas are often in close proximity to the umbilicus, ribs, or groin, which may interfere with pouch selection and adherence (6).
- Stomal complications
 - Bleeding
 - Hemorrhage during the immediate postoperative period is caused by inadequate hemostasis (4).
 - Trauma to stoma caused by improper fitting pouch. A wafer cut too close to the stoma can injure the delicate tissue. Stomal lacerations can occur as a result of the edge of the wafer rubbing back and forth against the side of the stoma (4).
 - Necrosis: Caused by ischemia and may be superficial or deep. Necrosis extending below the facial level may lead to perforation and peritonitis, requiring additional surgical intervention (4).
 - Mucocutaneous separation: This condition is caused by a breakdown of the suture line securing the stoma to the surrounding skin, leaving an open wound next to the stoma.
 - Prolapse: Telescoping of the bowel out through the stoma. In infants this condition is frequently related to poorly developed fascial support or excessive intra-abdominal pressure caused by crying.
 - Retraction: The stoma is flush or recessed below the skin surface. This condition may result from insufficient mobilization of the mesentery or excessive tension on the suture line at the fascial layer, excessive scar formation, or premature removal of a support device (4).
 - Stenosis: The lumen of the ostomy narrows at either the cutaneous level or the fascial level. Sudden decrease in output may indicate stenosis.
- Peristomal complications
 - Dermatitis
 - Allergic dermatitis
 - Contact dermatitis: Most common type of peristomal skin complication seen, generally from the leakage of fecal effluent on the skin.
 - Infection
 - Bacterial
 - Candidal
 - Mechanical trauma: Epidermal stripping, abrasive cleansing techniques, or friction due to illfitting equipment are the most common causes of mechanical injury to the peristomal skin.
 - Hernia: A peristomal hernia appears as a bulge around the stoma that occurs when loops of the bowel protrude through a facial defect around the stoma into the subcutaneous tissue (4).



FIG. 40.2. Immediately postoperative loop ileostomy. Segment of bowel on left is the exteriorized perforation from necrotizing enterocolitis.



FIG. 40.3. Premature infant with double-barrel colostomy.



FIG. 40.4. End ileostomy and wound closure with retention sutures posing a challenge for placing a pouch.

D. Ostomy Care

- Immediate postoperative care
 - Assess stoma for adequate perfusion.
 - Protect the stoma. Although stool output is usually minimal in the initial postoperative period, covering the stoma with an ostomy pouch prevents the mucosa from drying out, especially if the baby is under a radiant warmer. Petrolatum gauze may be used in the postoperative period if placement of the pouch has to be delayed because of problems with skin integrity.
 - Cover the mucus fistula with a moisture-retentive dressing to keep it from drying out. When securing a dressing on a neonate, use low-tack adhesives. There is increased risk of skin tears in neonates, especially when they are premature with delayed epidermal barrier development. Avoid placing petrolatum gauze over the pouching surface for the stoma, as it can impede adherence.
- Subsequent care
 - Regular assessment of the stoma
 - Protect peristomal skin from the effects of the effluent by pouching (Fig. 40.5). The effluent from a small bowel stoma contains proteolytic enzymes that can rapidly cause skin erosion. Ideally the bag should remain in place for at least 24 to 72 hours. The bag must be changed if there is any evidence of leaking effluent under the skin barrier wafer. Frequent bag changes, however, can result in denuded skin, especially in the premature infant (2,4,7). In situations with frequent leaking and pouch changes, expert help may be required to preserve the skin and obtain acceptable wear time. When a pouch cannot be maintained, it may be necessary to leave the pouch off and protect the peristomal skin with a protective barrier paste that will adhere to denuded skin to allow the skin to heal. The barrier ointment can be covered with petrolatum-impregnated gauze; fluff gauze can then be placed on top to absorb the effluent and changed as needed. In some cases of severe skin damage, some neonatal centers stop enteral feedings briefly to limit stool production and allow the skin to heal (2).
 - Protect stoma from trauma. Measures include accurate sizing of the opening to clear the stoma as the size changes. If the infant's movements cause the inner edge of the barrier to rub against the stoma, a moldable barrier between the stoma and the wafer can be used to protect the stoma.



FIG 40.5. One-piece ostomy appliance on small newborn dwarfs this infant, but provides longer wear time and holds larger volume of output than the preemie pouches previously used.

E. Equipment

Several different types of pouches and ostomy care supplies are available (Tables 40.2 and 40.3). One-piece pouches come with a barrier and pouch attached as a single unit. Two-piece appliances have a barrier and pouch separate, with a mechanism for attaching the pouch to the wafer, generally in the form of a plastic flange that snaps together. The type of pouch used with a neonate is generally either an open-end pouch that allows the passage of thick or formed effluent or a urostomy pouch with a spout designed for drainage of urine or liquid effluent. The type of pouch and the need for accessory products varies depending on the size of the child, the condition of the peristomal skin, abdominal size and contours, and institutional preference. In general, it is best to keep the procedure simple and to use as few products as possible (2). Special consideration needs to be given to the premature infant whose skin is immature and fragile. Several companies manufacture pouches for neonates and premature infants (Fig. 40.7). Neonatal units should have several varieties to choose from in order to meet each patient's individual needs.

Supplies

- Clean gloves
- Warm water
- Small basin or tub
- Clean, soft cloth
- 2 x 2-in gauze
- Appropriate-size pouch with clip/closure device
- Protective skin barrier wafer (if not using one-piece pouch)
- Other ostomy accessories as appropriate (Table 40.3 and Fig. 40.8).
- Scissors or seam ripper
- Stoma-measuring device

F. Applying the Pouch: Routine/Simple Ostomies (2,6,8)

- Remove old pouch by gently pulling up the edges and using water to loosen while pressing down gently on the skin close to the edge to reduce traction on the epidermis. Adhesive remover should not be used on a neonate <2 weeks of age. Limited use of adhesive remover, followed by thorough cleansing of the area to remove any chemical residue, is recommended only when the adhesive bond of the barrier to the skin is so strong that the skin might be injured during removal (2).
- Use damp soft gauze or paper washcloth to gently cleanse the stoma to remove adherent stool or mucus. It is common to have a little bleeding of the stoma when it is cleansed.
- Wash peristomal skin with water; pat dry. Soap is not recommended because it may leave a chemical residue that could cause dermatitis; furthermore, many soaps contain moisturizers that can adversely affect the adherence of the barrier to the skin. It is also not advisable to use commercial infant wipes, because most are lanolin-based and contain alcohol (2).
- Measure stoma(s) using stoma measuring device (Fig. 40.9). The opening generally is cut ¹/₁₆ to 1/8 in larger than the stoma, to limit the skin exposed to effluent. In tiny infants, in which the mucus fistula may be immediately adjacent to the functional stoma, one pouch may be sized to fit over both the stoma and the mucus fistula.

Туре	Advantages	Disadvantages
One piece (wafer	Simple and easy to apply	Difficult to customize, especially with
and pouch	Good choice for simple stomas	multiple stomas or abdominal irregularities
combined as one		Sometimes too large for small, premature
unit)		infants
Two piece (wafer	Easier to customize, especially for	More time-consuming to prepare
and pouch	multiple stomas, abdominal	More products to keep track of
separate)	irregularities	
Cut to fit with	Can be adjusted to meet changing	Must be sized and cut with each pouch
starter hole (may	stoma size, assuming starter hole is	change
be either one-piece	enot larger than stoma size	
or twopiece type		
pouch)		
Cut to fit without	Can be adjusted to meet changing	Must be sized and cut with each pouch
starter hole (may	stoma size	change
be either one-piece	eCan be customized to fit difficult	Difficult to start hole; must be careful not to
or two-piece type	stomas, multiple stomas, or to place	poke through bag with scissors
pouch)	stoma off-center in bag	
Precut hole (may	If stoma size is stable, less cutting	Stoma size may change and bag will not fit
be either onepiece	and adjusting needed with each bag	properly
or two-piece type	change	Difficult to stock large variety of custom
pouch)	Recommended for home use in older	sizes to meet needs of all NICU patients
	patients in whom stoma size will not	Not recommended for NICU stock
	change frequently	
Pouch choice of	Open end allows any type of effluent	Closure device may not hold for open-end
open-end,	to pass through. Urostomy allows	pouch. Tail of pouch is more difficult to
urostomy, or	liquid effluent to pass in a more	keep clean. Very thick liquid may not pass
closed-end	controlled manner. Closed end will	through the urostomy port, but can be
	not leak inadvertently from clip or	thinned by instilling water first. Closed-end
	closure failure.	pouch must be removed to empty.

TABLE 40.2 Ostomy Pouches

NICU, neonatal intensive care unit.

- Trace hole size onto wafer. Cut hole(s) using small scissors or a seam ripper (Fig. 40.10). After cutting, and before removing the paper backing, check the fit around the stoma and trim more if needed. Run a finger along the inside of the opening to make sure there are no sharp edges; these can be cut or smoothed by rubbing with the finger. It may be necessary to trim the wafer to avoid umbilicus, groin, etc. Cutting small slits along the edges of the wafer may help the barrier conform to the contour of the stomach.
- Warm wafer in hands to promote flexibility and enhance bonding to the skin. Avoid using a radiant heater to heat the wafer, because the amount of heat absorbed cannot be controlled and may burn immature skin (2).
- Press wafer to skin and hold for 1 to 2 minutes. Secure the edges of the wafer down to the skin to improve wear time. Avoid the use of high-tack adhesives. Pink tape is a waterproof tape that contains zinc oxide; it is very gentle and generally can be used safely. Other low-tack alternatives are paper tape, silicon tape, or clear film dressing.
- Change dressing to mucus fistula using a folded 2 x 2-in gauze piece and low-tack adhesive, or secure with diaper or tubular elastic dressing.

TABLE 40.3 Ostomy Accessory Products

Product Indications and Precautions

Barrier powder	This product is used to dry moist and/or weepy skin. It can add extra adhesiveness to the skin. It must be sealed by padding with a moistened finger and allowed to dry. In cases of severely moist
powaer	weeping skin, it may be necessary to apply powder and seal two or three times to attain a dry
	peristomal skin surface. It adds an additional barrier over the skin to protect from drainage.
	Apply in limited amounts and wipe off excess. Protect infant from inhalation of aerosolized
	powder by using minimal amounts and wiping away gently; do not blow powder away.
Paste	Barrier product that is semiliquid because of addition of alcohol. Best if applied to barrier and
	allow to air for 1 to 2 minutes to allow the alcohol to evaporate (Fig. 40.6). Not recommended
	for use on premature infants or term infants <2 weeks old.
Skin	Sealants use plasticizing agents to form a barrier on the skin that can protect from effluent and
sealants	also improve adherence of some adhesives. Most skin sealants contain alcohol and are therefore
	contraindicated for use in preemies or term neonates <2 weeks old. One skin sealant that does
	not contain alcohol is Cavilon No Sting Barrier Film (3M, St. Paul, MN, USA).
Moldable	Barriers that are adhesive and can be shaped to fill in uneven spaces; generally hold up very well
barrier	to corrosive effluent. Common types are Eakins Seals (Convatec, Princton, NJ, USA), and
	Barrier No. 54 (Nu-Hope Laboratories, Pacoma, CA, USA)
Caulking	Similar to moldable barriers, but come in narrow strips; they can be used to provide an extra
strips	barrier between the edge of the stoma and the barrier. May come in contact with stoma; soft
-	enough that it does not injure the mucosa. Examples are Ostomy Strip Paste (Coloplast, Marietta,
	GA, USA) and Skin Barrier Caulking Strips (Nu-Hope Laboratories, Pacoma, CA, USA).
Belt	Elastic belt with tabs that fit to ostomy pouch of some two-piece appliances. Belt can help
	maintain the appliance in place by holding it firmly to abdomen. Generally used as a last resort
	when unable to obtain acceptable wear time.



FIG 40.6. Barrier paste applied to wafer.



FIG 40.7. Examples of appliances for pouching a neonate.

G. Emptying the Pouch

- Supplies
 - Clean gloves
 - Diaper or syringe for withdrawing stool/effluent
 - o 30- to 60-mL syringe for irrigating/washing the bag
 - Tap water
- The pouch should be emptied when it is one third to one half full. Gas must also be released or vented to prevent pulling the adhesive wafer away from skin. Neonates generally produce large amounts of gas, related to increased intake with sucking and crying (2). Effluent can be drained directly into a diaper or withdrawn from the bag with a syringe. Use of two or three cotton balls placed in an open-end pouch can improve wear time by wicking the effluent away from the barrier and also may facilitate easy drainage of the pouch. It is generally not necessary to wash the pouch, but it may be necessary to add fluid to help loosen up thick or pasty stool. For the hospitalized neonate, measurement of ostomy output is usually indicated.
- Close the pouch with a closure device or rubber band.



FIG 40.8. Examples of ostomy accessories.

H. Complicated Stomas and Peristomal Skin Problems (5,9)

Table 40.4 lists complications and interventions for treating complex stomas and common stoma problems. Note that many of items used are not generally recommended for use on premature neonates or neonates <2 weeks of age, but in situations of deterioration of the peristomal skin they are sometimes used cautiously to prevent further deterioration and maintain an effective seal.



FIG 40.9. Measuring the stoma.



FIG 40.10. Cutting a hole in the wafer.

I. Vesicostomy Care

A vesicostomy does not require pouching; urine drains directly into the diaper. Care is similar to general perineal care of normal newborns (4). Occasionally, skin breakdown does occur; it can be treated with moisture barrier products and frequent diaper changes.

Gastrostomy Tubes

A. Indications For indications and insertion technique, see Chapter 39.

B. Types of Tubes See Table 40.5.

C. Gastrostomy Care

- Assessment
 - The health care provider must know if the patient has undergone a Nissen fundoplication or other antireflux procedure together with the gastrostomy.
 - Tolerance to feedings
 - Type and size of tube
 - Insertion site
 - Condition of the peristomal skin
- Special considerations for patients with Nissen or other antireflux procedure
 - Patient cannot vomit or burp.
 - Vent tube after crying and at first sign of gagging, discomfort, or distress.
- Gastrostomy tube site and routine skin care (6,10)
 - Clean gastrostomy tube site two to three times per day in the postoperative period and once per day after the site has healed. Use normal saline and sterile cotton swabs in the early postoperative period. Use mild soap and water after the site has healed. Diluted hydrogen peroxide (50% hydrogen peroxide and 50% water) is not recommended unless the site has dry, crusted blood (9).
 - Ensure that the antimigration device is flush against skin and the gastrostomy tube has not migrated.
 - Position tube at 90-degree angle.
 - A bottle nipple placed over the tube with the flanges resting on the abdominal wall may also be used to keep the tube at a 90-degree angle; secure with tape (Fig. 39.5).
 - Stabilize gastrostomy tube to prevent excess movement of tube, to decrease risk of stoma erosion, infection, bleeding, and development of granulation tissue.
 - Use an anchoring device (e.g., Hollister Tube Drainage Attachment Device, Hollister Inc., Libertyville, IL, USA) if the patient is allergic to tape or as a routine to secure the tube to skin.
 - Rotate bolster, flange of nipple, or wings of button every 4 to 8 hours to prevent pressure necrosis of skin. Do not place gauze between skin and bolster. A tension tab can be created by placing tape on the tube and pinning it to the diaper. A one-piece shirt with snap enclosure or tubular elastic dressing can also be used to cover the tube.
 - Assess site and peristomal skin for leaking, irritation, redness, rashes, or breakdown.
 Erythema and a minimal amount of clear drainage are to be expected in the first postoperative week.

D. Gastrostomy Tube Complications

Table 40.6 lists interventions for treating complications related to gastrostomy tubes.

	TABLE 40.4 Complications and Complex Ostomies
Complication	Interventions
Multiple stomas	Customize pouch to fit around or accommodate stomas in bag; mucus fistulas may
	be in or out of pouch.
Open incision or wound	Two-piece pouches without starter hole may allow for easier customization.
	Keep wound as clean as possible.
	Use hydrocolloid wound dressing (e.g., DuoDerm, Convatec; Replicare, Smith and
	Nephew, Largo, FL, USA) or calcium alginate in wound bed covered with a piece
	of clear film dressing to protect wound from stool.
	Paste and powders may also be used to protect peristomal skin.
	In some cases it may not be possible to apply a pouch; however, the skin must be
	protected from caustic effluent. A combination of a skin barrier such as Sensicare
	(Convatec) and petrolatum has been found to be effective.
Flush/retracted stoma	Apply paste or moldable barrier around hole in wafer.
	Use convex insert/convex pouch and belt to push skin back and allow stoma to
	protrude.
Prolapsed stoma	Notify surgeon if evidence of circulatory compromise.
	Protect the stoma from injury. When using twopiece pouch with plastic flange, the
	stoma could be pinched in the flange that secures the pouch to the wafer when
	closed.
	Adjust size of hole accordingly; cover exposed skin with moldable barrier or paste.
Peristomal hernia	Use a flexible wafer and pouching system to adjust to contour of the skin.
Mushroom-shaped stoma	aModify opening to accommodate size of "crownâ€; protect skin around base
	with moldable barrier or paste.
Irritant dermatitis	Ensure that hole is cut to fit properly.
	Use paste to protect from leakage.
	Apply powder to open, weepy skin.
	Assess for sensitivity to products.
	Apply topical steroids if needed to decrease inflammation, pain, and itching.
Peristomal Candida	Appears as red, shiny, macular, papular rash that is puritic.
albicans	Apply topical antifungal powder (e.g., Mycostatin) to skin. The powder should be
	mixed with a small amount of water, painted smoothly on the skin with a cotton
	swab, and allowed to dry before placing the appliance. Continue to use with each
	pouch change until rash resolves.
	Dry skin completely when changing pouch.
	Resize pouch so that no skin is exposed.
Dehydration, metabolic	Monitor intake and output carefully, especially for infants with ileostomy and/or
acidosis, electrolyte	high output.
imbalance	Assess lab values regularly. Infants can develop electrolyte imbalance rapidly.
Data from refs. (3,4,5,6,	7).
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Temporary/traditional Gastrostomy feeding tubes		Most commonly used as initial tube following Stamm procedure; long, self-retaining catheters of latex or silicone rubber with self-retaining devices (i.e., balloon)	Malecott (Bard, Covington, GA, USA) (collapsible wings), dePezzer (mushroom)		
		Silicone catheter with antimigration device and end cap	MIC (Ballard Medical, Draper, UT, USA), Corpak (Med		
Skin surface de	vices	Intended for use in established gastrostomy tract; have self-retaining devices, antimigration devices, and antireflux valves; two types, balloon and "Malecott type	Bard Button (Bard, Covington, GA, USA), Mic-key		
	TA	ABLE 40.6 Interventions for Gastrostomy Tube Co	omplications		
Complication	Inter	ventions			
Leaking at	Ensur	e that tube is properly situated in stomach; pull back	gently until resistance is met.		
Insertion site	Chaol	r water volume if holloon ture astheter			
Stollia	Ensur	water volume in balloon-type catheter.	cosal lining and skin		
cinargement	Use n	roper feeding attachment	cosar mining and skin.		
	Ensur	Ensure that tube is properly flushed and cleaned			
	Prote	Protect skin with skin barrier (e.g., Stomahesive wafer, paste or powder, Convatec; ILEX			
	paste,	paste, Convatec; or hydrocolloid dressing).			
	Use f	bam dressing (e.g., Hydrasorb, Convatec; Allevyn, Sr	nith and Nephew) rather than		
	gauze	gauze to wick moisture away from skin.			
	If not	If not contraindicated, consider H2-blocker and prokinetic agent.			
	Placin	Placing larger-size tube may temporarily control leakage but will not amend problem and is			
	Place	smaller tube and secure well to allow stoma to contra	act around the tube		
Dislodgement	Do no	t reinsert if <2 wks postop. Contact surgeon immedia	ately.		
2 1010 08011011	If >2	wks postop, replace as soon as possible (see Chapter	39).		
Bilious residual	sAsses	s for migration of tube (particularly if Foley is being	used).		
Bilious	Migra	tion results from inadequate stabilization. Tube may	migrate upward, causing		
vomiting	vomit	ing and potential aspiration, or downward, causing ga	astric outlet obstruction.		
Abdominal	Migra	ation into the small intestine can cause "dumping s	yndrome. When using a balloon		
distention	cathe	ter and migration is not recognized, inflation of the ba	lloon can lead to esophageal,		
D '	duode	enal, or small bowel perforation.			
Pain	Pull b	ack, if migrated, and secure.			
	Const	ube.			
Granulation	Norm	al finding: caused by proliferation of granulation epit	helial tissue in response to		
tissue	inflan	amation and irritation by foreign body	nenar ussue în response to		
	Preve	nt by stabilizing the tube.			
	Treat	by cauterizing with silver nitrate.			
	For la	rge amount of granulation tissue, consider applying T	Triamcinolone cream 0.5% 2-3		
	times	daily until resolved.			
Bleeding	Apply	gentle pressure to site.			
	Stabil	ize the tube.			
T	If gra	nulation tissue is present, treat appropriately.	1		
Irritant	Prote	ct skin with skin barrier (e.g., Stomahesive wafer, pas	te or powder, ILEX paste, or		
aermatitis	hydro	colloid dressing).			

TABLE 40.5 Types of Gastrostomy Tubes Description Examples

Туре

	Use foam dressing (e.g., Hydrasorb, Allevyn) rather than gauze to wick moisture away
	from skin.
	Assess for sensitivity to products/latex.
Candida	Apply topical antifungal to skin.
albicans	Control leakage.
	Dry skin completely after cleaning.
	Patient should also be assessed for oral thrush.
Clogged tube	Flush well after medications with 5 mL lukewarm water.
	A small amount (3-5 mL) of carbonated soda or cranberry juice may also be poured into the
	tube. Allow to set for 10 min, then flush with water.
Infection	G-tube site infections are uncommon; cellulitis is treated with systemic antibiotics.
Data from refs	(248911)

Data from refs. (2,4,8,9,11).

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References

1. Gauderer MWL. Stomas of the small and large intestine. In: O'Neil JA, Rowe MI, Grosfeld JL, et al., eds. Pediatric Surgery. 5th ed. St. Louis, MO: Mosby; 1998:1349–1359.

2. Rogers VE. Managing preemie stomas: more than just the pouch. J Wound Ostomy Continence Nurs. 2003;30:100–110.

3. Metcalfe P, Schwarz R. Bladder exstrophy: neonatal care and surgical approaches. Wound Ostomy Continence Nurs. 2004;31:284–292.

4. Erwin-Toth P, Doughty DB. Principles and procedures of stomal management. In: Hampton BG, Bryant RA, eds. Ostomies and Continent Diversions: Nursing Management. St. Louis, MO: Mosby; 1992:29–104.

5. Craven DP, Fowler JS, Foster ME. Management of a neonate with necrotizing enterocolitis and eight prolapsed stomas in a dehisced wound. J Wound Ostomy Continence Nurs. 1999; 26:214–220.

6. Borokowski S. Pediatric stomas, tubes, and appliances. Pediatr Clin North Am. 1998;45:1419–1435.

7. Garvin G. Caring for children with ostomies and wounds. In: Wise B, McKenna C, Garvin G, et al., eds.

Nursing Care of the General Pediatric Surgical Patient. Gaithersburg, MD: Aspen; 2000:261–278. 8. Borkowski S. Gastrostomy tube stabilization and security. Sutureline. 2005;13:8–10.

9. Association of Women's Health, Obstetric and Neonatal Nurses, National Association of Neonatal Nurses. Evidence-Based Clinical Practice Guideline: Neonatal Skin Care. Washington, DC: AWHONN; 2001.

10. Borkowski S. Gastrostomy surgery and tubes. Sutureline. 2000;8:1-3.

11. Colwell JC. A practical guide for the management of pediatric gastrostomy tubes based on 14 years of experience. J Wound Ostomy Continence Nurs. 2004;31:193-200.

41-Transfusion of Blood and Blood Products

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Overview

Blood Products Utilized in Neonates

- Packed red blood cells (PRBCs)
- Whole or reconstituted whole blood
- Platelet concentrates derived from whole blood or plateletpheresis
- Fresh frozen plasma (FFP) or frozen thawed plasma
- Cryoprecipitate
- Granulocyte concentrates derived from granulocytopheresis

Sources of Blood Products

- Banked donor blood
- Directed donor transfusions
- Autologous fetal blood transfusions

Indications, requirements, and transfusion techniques differ for each procedure and component. Simple transfusions are discussed in this chapter. Exchange transfusions are discussed in Chapter 42. Complications common to all blood products are listed later in this chapter.

A. Precautions

- Whenever possible, obtain informed consent prior to transfusions, delineating risks, benefits, and alternatives to transfusion.
- Limit use of transfusions to justified indications.
- Select blood product appropriate for infant's condition.
- Confirm with proper identifiers at bedside that blood product is for correct patient. Maintain all records relevant to collection, preparation, transfusion, and clinical outcome.
- Avoid excessive transfusion volume or rate unless acute blood loss or shock dictates faster transfusion.
- Store blood and blood products appropriately. Freezing and lysis may occur if red blood cells (RBCs) are stored in unmonitored refrigerators.
 - Use blood bank refrigerator for storage of RBCs, whole blood, thawed FFP, and thawed cryoprecipitate.
 - Temperature should be controlled at $4 \hat{A} \pm 2\hat{A}^{\circ}C$ with constant temperature monitors and alarm systems.
 - Refrigerator should be quality-controlled at least daily.
 - Designated for blood products only
 - $_{\odot}$ Store platelets at 20 to 24Å $^{\circ}C$ with continuous agitation.
- PRBCs and whole blood should be out of refrigeration for <4 hours to minimize bacterial contamination and red blood cell hemolysis.
- Use approved blood warming services for PRBCs and whole blood. Syringes for aliquots must not be warmed in water baths because of the risk of contamination.
- Stop transfusion or slow rate if baby manifests any adverse side effects.
 - o Tachycardia, bradycardia, or arrhythmia
 - Tachypnea
 - Systolic blood pressure increases of >15 mm Hg, unless this is the desired effect

- \circ Temperature above $38\hat{A}^{\circ}C$
- Hyperglycemia or hypoglycemia
- Cyanosis
- Skin rash, hives, or flushing
- Hematuria/hemoglobinemia
- Hyperkalemia
- Transfuse RBCs cautiously in incipient or existing cardiac failure.
 - Monitor heart rate, blood pressure, and peripheral perfusion.
 - Transfuse PRBCs, or
 - Consider partial exchange transfusion:
 - With hemoglobin level <5 to 7 g/dL
 - With cord hemoglobin <10 g/dL
 - Prevent fluctuations in glucose during RBC transfusion:
 - In infants weighing <1,200 g or in other unstable infants, to prevent hypoglycemia
 - Do not discontinue parenteral glucose administration.
 - Establish separate intravenous line for blood administration.
 - When transfused blood has elevated glucose concentration, expect rebound hypoglycemia in infants with hyperinsulinism.
- B. Pretransfusion Testing and Processing (1, 2)
 - Blood group and Rh type
 - Maternal ABO blood group and Rh type: Screen maternal serum for atypical antibodies.
 - Baby's ABO blood group and Rh type: Screen baby's serum for atypical antibodies if maternal blood is unavailable.
 - Cord blood may be used for initial testing.
 - Baby's blood group is determined from the red cells alone, because the corresponding isoagglutinins anti-A and anti-B are usually weak or absent in the serum.
 - Cross-matching
 - Compatible blood may be low-anti-A, anti-B titer Group O Rh-negative blood or blood of the infant's ABO group and Rh type (except in alloimmune hemolytic disease of the newborn).
 - Conventional cross-match is not required if infant <4 months old and no atypical antibodies are detected.
 - Compatibility testing for repeated small-volume transfusions is usually unnecessary, because formation of alloantibodies is extremely rare in the first 4 months of life.
 - If antibody screen is indirect antiglobulin test (IAT)-positive in mother or baby:
 - Serologic investigation to identify antibody(ies) is necessary.
 - Full compatibility testing is required.
 - If anti-A or anti-B detected in infant, infant should receive RBC lacking A or B antigen until antibody screen is negative.
 - If infant has received large volumes of plasma or platelets, passive acquisition of antibodies may occur; cross-matching is recommended.
 - If directed donor blood from a parent is used, cross-matching is required.
 - Specially processed products
 - Blood donated in the United States is tested for human immunodeficiency virus (HIV), human T-lymphotropic virus I/II, hepatitis B and C viruses, West Nile virus, syphilis, and bacteria (in platelet products).
 - Cytomegalovirus (CMV)-seronegative or third-generation leukodepleted (LD) blood is recommended for infants with birthweight 1,200 g born to seronegative mothers or those with unknown serostatus (1).
 - Use of universal LD and/or CMV seronegative products is institution-specific (3).
 - Irradiation to prevent transfusion-associated graft-versus-host disease (TAGVHD)

- Whole blood, PRBCs, previously frozen RBCs, granulocyte and platelet concentrates, and fresh plasma have been implicated in TAGVHD; LD products have also been implicated.
 - Clinical indications for irradiated blood components are (1, 2, 4, 5 and 6)
 - Intrauterine transfusion
 - Premature infants, variably defined by weight and postgestational age
 - Congenital immunodeficiency suspected or confirmed
 - Undergoing exchange transfusion for erythroblastosis
 - Hematologic/solid organ malignancy
 - Recipient of familial blood donation
 - Recipient of HLA-matched product
- Some institutions provide irradiated blood products to all neonates to avoid TAGVHD in patients with undiagnosed immunodeficiency.

C. Equipment

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- Blood product (see Appendix C)
- Cardiorespiratory monitor
- Blood: All blood and blood components must be filtered immediately prior to transfusion despite prestorage LD. Many transfusion services supply PRBCs and occasionally platelets and cryoprecipitate prefiltered to the neonatal intensive care unit (NICU).
 - $^{\circ}$ Administration set with in-line filter of 120- to 170-µm pore size to be used for all products
 - $_{\odot}$ $\,$ Microaggregate filter, 20- to 40- $\hat{A}\mu m$ pore size
 - Must follow manufacturer's instructions
 - Some function only if product is dripped
 - Not advisable for syringe administration
 - Usefulness questionable and unnecessary if LD and/or additive RBC's used
 - Prestorage LD filtration (2, 3, 6)
 - Removes 99.9% of white blood cells (WBCs) or 3 log leukodepletion
 - Must follow manufacturer's instructions
 - Attenuation/abrogation of CMV and other viruses (Epstein–Barr, HTLV I/II Human Thymphocyte Virus) harbored in WBCs
 - LD performed by collecting facility preferred to bedside use
- Sterile syringe
- Blood administration set
 - $_{\odot}~$ a May be manufactured with integral 120- to 170-ŵm filter
 - May not be needed if prefiltered product is dispensed from transfusion service
 - Blood warmer not needed
- Automated syringe pump with appropriate tubing and needle (7, 8, 9, 10, 11 and 12)
 - Least hemolysis occurs with straight syringe pumps
 - Vascular access: PRBCs may be transfused through 24-, 25-, or 27-gauge needles and short catheters but not through 27- or 28-gauge central venous catheters.
 - The amount of hemolysis that results from infusion of RBCs is directly proportionate to the age of the blood and the rate of transfusion and inversely proportional to needle size.
 - Hyperkalemia, hemoglobinuria, and renal dysfunction may result if hemolyzed blood is transfused.
- Normal saline flush (1 mL or more) to clear intravenous solution from line.

Packed Red Blood Cell Transfusions

A. Indications

- Guidelines and justifications for transfusions are controversial and vary among different authorities and institutions (13, 14). The transfusion guidelines followed by the U.S. Multicenter Recombinant Human Erythropoietin Trial established the safety, at least in the short term, of conservative transfusion practices for infants needing mild and moderate respiratory support (Table 41.1) (15). Transfusion guidelines continue to be liberal for infants with severe cardiorespiratory compromise (4). Limiting transfusion is now questioned as a cause of central nervous system complications and apnea (16).
- In general:
 - Replace acute blood loss.
 - Correct anemia that is compromising cardiovascular status or oxygen-carrying capacity.

B. Contraindications

- None absolute
- Exert caution in patient with:
 - Volume overload
 - Congestive heart failure
 - T-activation

TABLE 41.1 Guidelines for Transfusion of RBCs in Patients <4 Months of Age</th>

- 1. Hematocrit < 20% with low reticulocyte count and symptoms of anemia^a
- 2. Hematocrit <30% with an infant:
 - a. On >35% hood O_2
 - b. On O_2 by nasal cannula
 - c. On continuous positive airway pressure and/or intermittent mandatory ventilation with mechanical ventilation with mean airway pressure <6 cm H_2O
 - d. With significant apnea or bradycardia^b
 - e. With significant tachycardia or tachypnea^c
 - f. With low weight gain^d
- 3. Hematocrit <35% with an infant:
 - a. On >35% hood O_2
 - b. On continuous positive airway pressure/intermittent mandatory ventilation with mean airway pressure 6 cm $\rm H_2O$
- 4. Hematocrit <45% with an infant:
 - a. On extracorporeal membrane oxygenation
 - b. With congenital cyanotic heart disease

^aTachycardia, tachypnea, poor feeding.

^bMore than six episodes in 12 hours or two episodes in 24 hours requiring bagand-mask ventilation while receiving therapeutic doses of methylxanthines.

^cHeart rate >180 beats/min for 24 hours, respiratory rate >80 breaths/min for 24 hours.

^dGain of <10 g/day observed over 4 days while receiving 100 kcal/kg/day.

Source: Adapted from **Roseff SD**, **Luban NL**, **Manno CS**. Guidelines for assessing appropriateness of pediatric transfusion. *Transfusion*. 2002;42:1398-1413.
C. Technique

- Determine total amount of blood needed.
 - Calculate volume of blood for transfusion. Most infants are transfused 10 to 15 mL/kg of PRBCs, which will increase the hemoglobin by 3 g/dL.
 - Calculate volume of PRBCs required:

[EBV — (Hct desired - Hct observed)] Hct of PRBC unit

where EBV is the estimated patient's blood volume, approximately 80 to 85 mL/kg in full-term infants and approximately 100 to 120 mL/kg in preterm infants, and Hct is hematocrit.

- Include volume of blood needed for dead space of tubing, filter, pump mechanism (varies from system to system; may be as much as 30 mL).
- Obtain blood product (see Appendix C).
 - Several studies have documented the safety of using PRBCs stored in additive anticoagulant preservative solutions to outdate.
 - Avoid use of RBCs stored in additive-containing solutions for massive transfusions, unless the additive is removed by inverted storage or centrifugation; risks of hyperosmolality, hyperglycemia, hypernatremia, hyperkalemia, hyperphosphatemia are postulated (17).
 - In infants receiving 10 to 15 mL of PRBCs, the amount of K^+ delivered by the transfusion is estimated to be only 0.15 mEq/kg and does not pose a significant risk to most neonates.
 - Donor exposures are reduced by repeatedly obtaining small aliquots of PRBCs from dedicated units until the outdate of 35 to 42 days (Figs. 41.1 and 41.2) (18, 19).



FIG. 41.1. Neonatal syringe set with filter. (Courtesy of Charter Medical Ltd., Winston-Salem, NC, USA). This system, when used with sterile connection technology, provides a closed delivery system that maintains primary unit outdate. Syringe blood aliquots (PRCBs, plasma) must be administered to the patient within 24 hours and syringe platelet aliquots within 4 hours.

- Verify whether cross-matched product is necessary or un–cross-matched product is adequate.
- Confirm that restrictions have been adhered to on blood product and transfusion tag.
 - CMV: Tested/untested
 - Irradiated: Yes/no
 - Directed (familial) donation: Yes/no
 - RBC antigen-negative: Yes/no
 - Sickle tested-negative: Yes/no
 - Other restrictions specified: Yes/no
- Verify appropriateness of blood selected for patient by comparing blood product and unit tag (integral to blood unit) information and patient identification. Bar-code reading devices are advisable.
 - Blood unit tag and blood bag
 - Patient hospital or medical record number
 - Patient identification by armband or footband
 - Blood group and type of both donor and recipient
 - Expiration date and time and restrictions on unit and order
 - Restrictions as ordered by physician or by institutional guidelines
- Warming PRBCs (2, 20, 21)
 - There is no need to warm small-volume PRBC aliquots, particularly if the transfusion is given over 2 to 3 hours.
 - PRBCs may be warmed by placing the syringe beside the infant in the warm-air incubator for 30 minutes prior to transfusion.
 - Inappropriate warming by exposure of blood to heat lamps or phototherapy lights may produce hemolysis.
- Adhere to sterile technique throughout procedure.
- If prefiltered PRBCs are provided by the blood bank in a syringe, attach tubing directly to syringe.
- If PRBCs are provided in a bag, use large-bore needle (18-gauge or larger) to withdraw volume into syringe. Filter should be placed between bag and syringe (Fig. 41.1).
- Prime tubing with blood. Clear syringe and tubing of bubbles, and mount into infusion device.
- Verify patency of vascular access.
- Clear line into patient with normal saline.
- Record and monitor vital signs.
- Determine spot glucose test. Repeat hourly as needed.
- Begin transfusion at controlled rate: 5 mL/kg/hr.
- Gently invert container of blood every 15 to 30 minutes to minimize sedimentation.
- Stop transfusion if any adverse change in condition occurs.
- At end of infusion, clear blood from line with saline.
- Check recipient hemoglobin and hematocrit, if necessary, at least 2 hours after transfusion.
- If posttransfusion hematocrit/hemoglobin is not up to expected level, consider:





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FIG. 41.2. Use of a sterile connecting device. A: An adult RBC unit is shown along with a set of pediatric transfer bags. The transfer bags can be attached by spiking the unit, causing it to expire in 24 hours; alternatively, the transfer bags can be connected using a sterile connection device. B, C: The separate tubings are loaded into the tubing holders of the device. The covers are closed. D: A welding wafer heated to about $500\hat{A}^\circ$ F melts through the tubing. The tubing holders realign and the welding wafer retracts allowing the tubing ends to fuse together. E: The unit can now be aliquoted as needed. Because a functionally closed system has been maintained, the expiration date of the blood has not changed.

- Low hematocrit of PRBC unit
- Inappropriate calculation of transfusion requirement
- Ongoing blood loss
- Transfusion reaction
- \circ $\;$ Hemolysis due to ABO or other RBC incompatibility $\;$
 - Infant has circulating anti-A, anti-B, and anti-AB, which is bound to A or B antigens on transfused RBCs.
 - Direct antiglobulin test (DAT) negative initially but now positive

- Unexpected increase in bilirubin
- Infant has RBC antibody other than ABO.
- Hemolysis from extrinsic damage (mechanical) to RBCs or donor has hemolytic disorder.

Whole or Reconstituted Whole Blood

Reconstituted whole blood is prepared by adding a unit of RBCs to a compatible unit of FFP (1, 2). Levels of coagulation factors V and VIII are low, platelet function is grossly abnormal, and concentration of K^+ is high in stored whole blood. Transfusion of PRBCs and other blood components as required is preferable to the use of stored whole blood.

A. Indications

- Massive transfusion as in acute blood loss, when restoration of blood volume and oxygen-carrying capacity are needed simultaneously.
- Exchange transfusions
- Cardiopulmonary bypass
- Extracorporeal membrane oxygenation
- Continuous hemofiltration
- In children <2 years of age, transfusion of fresh (<48 hours old) whole blood has been associated with significantly less postoperative blood loss following bypass surgery for complex congenital heart disease (22). Recent studies now question use of whole blood (23).

	TABLE 41.2 Guidelines for Platelet Transfusions (× 10 ⁹ /L)				
	Nonbleeding, Sick, Nonbleeding		Nonbleeding,	Before Invasive Actual	
	Preterm	Stable, Preterm	Term	Procedure	Bleeding
Roberts (2006) Canada	<50	<30	<30	<50	<100
Roseff (2002) USA	<100	<50	<30	<50 if failure of products	<50 if stable <50 if sick
Gibson (2004) UK	<30	<20	<20	<100 if DIC Not specified	<50

 $\hat{a}\in \infty$ Sick $\hat{a}\in$ infants in this context defined as those with a history of perinatal asphyxia, extremely low birthweight (<1,000g), need for ventilatory support with an inspired oxygen concentration >40%, clinically unstable, signs of sepsis, or those who require numerous invasive interventions.

B. Precautions

- Not suitable for simple transfusion for anemia
- Not suitable for correction of coagulation factor deficiencies
- Hyperkalemia may result from rapid transfusion of large volumes (10, 11, 24).
- Anticoagulant effects must be considered (17).

C. Equipment and Technique

- Same as for PRBCs
- The rate of transfusion may be increased to 10 to 20 mL/kg/hr to replace acute blood loss.
- When used for complex mechanical procedure, 120- to 150-µm filters often in line to device.

Platelet Transfusions

A. Indications (Table 41.2)

The platelet count at which transfusion is recommended has to be individualized, because hemostatic competence is determined not only by the quantity of platelets but also by platelet function, vascular integrity, levels of coagulation factors, and underlying disorder/disease.

B. Contraindications

Because the half-life of platelets is reduced in conditions of rapid consumption and/or immune destruction, platelet transfusions often do not produce a measurable platelet increment. Their use in these conditions is reserved for life-threatening situations in which other therapy is ineffective. In infants previously exposed to heparin, who develop thrombocytopenia, and platelet refractoriness, consider heparin-induced thrombocytopenia as cause (25). Poor posttransfusion increment in infants with neonatal alloimmune thrombocytopenia (NAIT) can define diagnosis (26).

C. Equipment and Technique

- Platelets
 - Random donor platelet concentrate (5.5 3 \tilde{A} 10¹⁰ platelets in 40 to 70 mL of plasma)
 - Separated from whole blood by centrifugation within 8 hours of blood draw
 - Resuspended in plasma
 - Shelf life of 5 days
 - Volume-reduced platelets
 - Standard platelet concentrate concentrated to a volume of 15 to 20 mL by centrifugation (27)
 - Associated with loss of platelets and possible decrease in platelet function
 - Shelf life reduced to 4 hours
 - Use only if infant has oliguria, severe volume load sensitivity
 - Plateletpheresis (3 \tilde{A} 10¹¹ platelets in volume of 250 mL plasma)
 - Removes only platelets, returns RBCs and plasma to donor
 - Usually leukodepleted before storage
 - Permits repeated donations from same donor every 48 hours under select circumstances
 - High yield of platelets
 - More expensive product
 - Useful when multiple platelet transfusions of a particular antigen specificity are required, as in neonatal albimmune thrombocytopenia (NAIT) or for infants on extracorporeal membrane oxygenation (ECMO) needing multiple platelet transfusions.
 - May be HLA-typed or typed for PLA1 or other specific platelet antigen in case of NAIT.
 - Maternal plateletpheresis product is preferred for NAIT. Use maternal antigennegative platelets, washed, irradiated, and resuspended in group-compatible ABO plasma or saline.
- Calculate volume of platelets to transfuse based on type of product.
 - 10-15 mL/kg of random platelet provides $10 \text{ }\tilde{A}$ 10^9 platelets/kg and should increase platelet count to >50 \tilde{A} 10^9 /L in the absence of ongoing consumption.
 - Can use same calculations for apheresis platelets, but studies do not confirm posttransfusion increments. Advise use of equivalent unit calculations and not mL/kg.
 - Other products (HLA-typed, cross-matched platelets) used for platelet (PLT) refractoriness; washed PLTs if using PLT antigen-matched maternal PLTs.

- Blood administration set with 120- to 170- $\hat{A}\mu m$ in-line filter, unless platelets have been prefiltered while drawing into a syringe. Specific sets designed for plasma/ platelets have in-line filters with reduced surface area to increase platelet transfusion efficacy.
- Sterile syringe for automated pump infusion. Use of syringe technique will increase damage to platelets. Administer by drip if clinically feasible.
- Automated syringe pump
- Connecting intravenous tubing
- Intravenous access, preferably through 23-gauge or larger needle or through umbilical venous catheter
- Normal saline flush solution

D. Precautions

- Use type-specific (Rh-negative) platelets when potential for sensitization is present (i.e., in Rhnegative female). Although platelets do not have Rh antigens, all products have some red blood cell contamination (less in plateletpheresis), which may cause Rh sensitization (28).
- Use platelets from donor with ABO-compatible plasma (Table 41.3). Isohemagglutinins in ABO-incompatible plasma may result in hemolysis, a positive DAT, and poorer in vivo platelet survival than anticipated.
- Transfuse platelets as soon after preparation as possible. Platelets should never be refrigerator-stored or warmed.
- Platelets should not be infused through arterial lines.

E. Technique for Platelet Administration by Automated Syringe

- Estimate by weight the volume of platelets in a single bag to determine fluid load to infant.
- Confirm correct platelet product.
 - Infant and unit identification

TABLE 41.3 Choice of Platelets

BabyDoorOOAA, ABBB, ABAB

- Infant and donor blood group, and Rh type
- Check other restrictions: CMV negative, irradiated, etc.
- Attach, aseptically, in sequence:
 - Platelet concentrate or bag aliquot
 - Platelet administration set, including filter
 - Three-way stopcock
 - Transfusion syringe
- Draw into syringe volume of platelets for transfusion and tubing dead space. Clear air bubbles.
- Remove syringe from three-way stopcock and attach to connecting tubing.
- Establish IV access. If infant is at risk for hypoglycemia with interruption of continuous glucose source, start new IV or monitor closely throughout infusion.
- Clear IV of glucose solution with 1 mL or more of normal saline.
- Attach connecting tubing and syringe to IV line.
- Monitor patient's vital signs.
- Infuse platelets over 1- to 2-hour period, faster if tolerated by infant.

- After infusion is complete, flush IV line with 1 mL of normal saline before restarting glucose solution.
- Determine survival time of transfused platelets by obtaining platelet counts at 1, 2, or 24 hours.

F. Complications

See list of complications for all blood products (1, 2, 29).

- Accentuated hemolysis in sensitized but IAT-negative ABO setup
- Rh sensitization in Rh-negative recipient (28)
- Volume overload
- Allergic reactions, including hypotension
- Transfusion-related lung injury (TRALI) (30, 31)
- Increased morbidity in necrotizing enterocolitis (NEC) (32)

Granulocyte Transfusions

A. Indications

- Granulocyte transfusions are used infrequently, displaced by improved cidal antibiotics, supportive care, and, in rare circumstances, use of granulocyte- and granulocyte/macrophage-stimulating factors (33).
- Historical indications not confirmed in randomized trials to be relevant:
 - Neonates <14 days old, with bacterial sepsis and neutropenia (neutrophil plus band count <3 $-10^{9}/L$)
 - $\circ~$ Older neonates with bacterial sepsis, unresponsive to antibiotics with neutrophil and band counts <0.5 10⁹/L

B. Equipment and Technique

- Granulocyte concentrates for neonatal use should be prepared by automated granulocytapheresis and contain 1 to 2 10⁹ neutrophils in 10- to 15-mL/kg volume. Steroid- or G-CSF-mobilized donor preferred.
- Daily transfusions may be necessary until there is clinical improvement and evidence of recovery of neutrophil counts.
- Component must be ABO- and Rh-compatible with recipient, and cross-match-compatible.
- Product should be irradiated, CMV-negative, and infused as soon as possible after collection.
- The product should not be refrigerated or warmed above room temperature.
- Standard 120- to 170-µm filters should be used for infusion; microaggregate and leukodepletion filters must be avoided.

C. Precautions

Storage of product for >8 hours is associated with a rapid decrease in WBC function, making this a less than useful product. Fever, alloimmunization, TRALI, and CMV have all been reported complications. Fresh Frozen Plasma, Frozen Thawed Plasma, and Cryoprecipitate

A. Indications (2, 34)

Clinically significant bleeding or for correction of hemostatic defects prior to invasive procedures in the presence of

- Complex factor deficiency unresponsive to Vitamin K
- Isolated congenital factor deficiency for which virus-inactivated-plasma-derived or recombinant factor concentrates are unavailable
- Congenital or acquired dys- or hypofibrinogenemia (cryoprecipitate is preferred)

B. Contraindications

- None absolute
- Exert caution when possibility of volume overload exists.
- Necrotizing enterocolitis (NEC): Use with caution. May aggravate hemolysis in infant with NEC and T-activation (35, 36).
- Not indicated for hypovolemic shock in the absence of bleeding, nutritional support, treatment of immunodeficiency, or prevention of intraventricular hemorrhage.

C. Equipment and Technique

See Platelet Transfusion.

- Cross matching is not required because type-specific or AB-negative product is usually issued.
- Dose of FFP is 10 to 20 mL/kg; multiple transfusions may be required until the underlying condition resolves.
- Once thawed, FFP should be transfused within 6 hours for labile factor replacement.
- In cases for which repeated FFP transfusions are required, a thawed unit from a single donor may be divided into smaller aliquots and used within 24 hours if stored between 1 and 6ŰC.

Directed Donor Transfusions

A. Potential Problems

Directed donations provide no known benefit in terms of increased safety and may have several disadvantages (37):

- Possible increased risk of transmitting infectious disease because directed donors are often first-time or infrequent donors with no track record of safety, unlike established volunteer donors, whose screening tests are negative repeatedly.
- Possibility of serologic incompatibility between the recipient baby and the family donors.
 - Plasma of biologic mothers may contain alloantibodies to blood cell antigens inherited by the infant from the father, which may lead to hemolysis, thrombocytopenia, and WBC aggregation in the infant, when transfused without the protective barrier of the placenta.
 - Paternal blood cells may express antigens to which the neonate may have been passively immunized by transplacental transfer of maternal antibodies.
 - Routine pretransfusion testing may not detect these serologic incompatibilities, which may result in hemolytic or other transfusion reactions.
- Although biologic parents may be interested in donating for their infants, many are likely to be ineligible for medical or serologic reasons.

B. Precautions

- Directed donations must be screened as stringently as volunteer donations.
- Biologic mothers should not provide blood components containing plasma; maternal red cells and platelets should be given as washed concentrates.
- Fathers and paternal blood relatives should preferably not serve as donors for blood components containing cellular elements; a full antiglobulin cross-match should be performed to detect red cell incompatibilities if their use is unavoidable.
- All blood components obtained from first- or second-degree relatives should be irradiated prior to transfusion.

Autologous Fetal Blood Transfusions (38,39,40,41)

A. Indications

- Delivery room resuscitation of infants with shock and profound anemia, when O Rh-negative bank blood is not readily available.
- Source of autologous PRBCs for elective transfusion to preterm infants.
- Source of cord blood for freezing for hematopoietic reconstitution.

Protocols for proper collection with appropriate anticoagulation, without bacterial contamination, are still being refined for these indications.

- **B.** Contraindications
 - Maternal infection
 - Chorioamnionitis
 - o Sepsis
 - Hepatitis, HIV
 - Prolonged rupture of membranes >24 hours

C. Complications

- Bacterial sepsis from contaminated collection
- Insufficient volume collected for transfusion
- Over/undercollection for volume of anticoagulant used

Complications of Blood Transfusions

Transfusions are safer now than ever before, but they are not risk-free (4, 5).

- Transmission of infectious diseases (42, 43)
 - Viruses: Risk varies geographically
 - Human immunodeficiency virus: Estimated potential risk in United States from a blood donor with negative serologic tests is <1 in 2,135,800.
 - Human T-lymphotropic virus I and II: Risk 1 in 641,000
 - Hepatitis B virus: Risk 1 in 1,205,000
 - Hepatitis C virus: Risk 1 in 1,935,000
 - Hepatitis A virus: Risk 1 in 1,000,000, asymptomatic in newborn, but may cause symptomatic infection in adults who are in contact with infected neonates.

- Cytomegalovirus: Transmitted by cellular blood products, not transmitted by FFP or cryoprecipitate; risk of transfusion transmitted CMV from cellular blood components from CMV-seropositive donors 8% to 25%, risk of CMV infection after transfusion of CMV-seronegative or effectively leukoreduced components is <1% to 4%.
- Hepatitis G, parvovirus B-19, Epstein–Barr virus
- Bacteria (44)
 - Persistent problem, often unrecognized
 - Platelet concentrates and PRBCs most often implicated
 - Organisms: Yersinia enterocolitica, Serratia, and Pseudomonas genus for RBCs; Staphylococcus epidermidis, Bacillus species for platelets.
 - Treponema pallidum: Does not survive refrigeration; syphilis reported after transfusion of fresh blood.
 - Red cell units with gross bacterial contamination may occasionally be identified by the darker color of the blood in the bag compared with the color of blood in the tubing.
- o Protozoan
 - Malaria: Rare in the United States but reported even in nonendemic areas (45)
 - Babesiosis
 - Chagas disease
- Creutzfeldt–Jacob
 - Few proven cases of transfusion-transmitted new-variant Creutzfeldt–Jacob disease at present (46)
 - Most blood collection centers attempt to minimize the risk by excluding donors considered to be at higher risk for possibly harboring the infection, by family and travel history and specific medical history (2).
- Hemolytic reactions
 - Acute hemolytic immunologic reactions: Rare, because of absence in infant of naturally occurring anti-A or anti-B antibodies, and infrequent posttransfusion red cell alloimmunization despite multiple transfusions.
 - T-activation: A severe form of immune-mediated hemolysis is associated with the transfusion of adult blood containing anti-T antibodies in neonates with activation of the normally cryptic T-antigen on the surface of neonatal red cell membranes (35, 36).
 - Common in premature infants and associated with necrotizing enterocolitis and sepsis
 - Suspect T-activation in neonates at risk with intravascular hemolysis, hemoglobinuria, hemoglobinemia following transfusion of blood products, or unexpected failure to achieve posttransfusion hemoglobin increment.
 - Routine cross-matching will not detect T-activation when monoclonal antiserum is used.
 - Diagnosis: By minor cross-matching, discrepancies in forward and reverse blood grouping, confirmed by specific agglutination tests using peanut lectins.
 - Use washed red cells, platelets, and low-titer anti-T plasma only when hemolysis is confirmed (35).
- Nonimmunologic causes of hemolysis
 - $\circ~$ Mechanical, through excessive infusion pressure through small needles or 20- to 40-Å μm filters
 - Accidental overheating or freezing of blood
 - Simultaneous administration of incompatible drugs and fluids
 - Transfusion of abnormal donor cells (glucose 6-phosphate dehydrogenase deficiency, hereditary spherocytosis)
- Other immunologic reactions
 - Transfusion-associated graft-versus-host disease

See processing with irradiation for risk factors and prevention.

- Transfusion-related acute lung injury
 - Secondary to transfusion of donor blood containing antigranulocyte and anti-HLA antibodies, leukoagglutinins directed against recipient granulocyte antigens, or to transfusion of plasma cytokines/mediators that affect recipient granulocytes, causing pulmonary endothelial adhesion defects
 - Initiates life-threatening hypoxemia secondary to complement activation and microvascular lung injury
 - Rarely reported in neonates; may be difficult to distinguish from other causes of respiratory deterioration in sick infants.
- Circulatory overload

Alterations in pulmonary compliance and blood pressure

- Adverse metabolic effects
 - Hyperkalemia
 - Blood that is irradiated and then refrigerator-stored may have K⁺ levels of 30 to 50 mEq/L or higher in the supernatant plasma.
 - Small-volume transfusions of stored red cells do not cause clinically significant elevations in serum K⁺ levels.
 - Life-threatening hyperkalemia has been described in sick infants and in those receiving rapid infusions of large volumes of stored red cells.
 - Washed or fresh (<14 days) red cells are recommended for infants with profound hyperkalemia, renal failure, or when large volumes are transfused rapidly.
 - Hypoglycemia or hyperglycemia
 - Hypocalcemia
 - Alterations in acid–base balance with large transfusions

References

1. American Association of Blood Banks. Standards for Blood Banks and Transfusion Services. 23rd ed. Bethesda, MD: AABB; 2004.

2. American Association of Blood Banks. Technical Manual. 14th ed. Bethesda, MD: AABB; 2003.

3. Ferguson D, Hebert PC, Lee SK, et al. Clinical outcomes following institution of universal leukoreduction of blood transfusions of premature infants. JAMA. 2003;289:1950–1956.

- 4. Luban NL. Neonatal red blood cell transfusions. Vox Sang. 2004;87(suppl 2):184–188.
- 5. Ramasethu J, Luban NL. Red cell transfusions in the newborn. Semin Neonatol. 1999;4:5–16.

6. Strauss RG. Data driven blood banking practices for neonatal RBC transfusions. Transfusion. 2000;40:1528–1540.

7. Frey B, Eber S, Weiss M. Changes in red blood cell integrity related to infusion pumps: a comparison of three different pump mechanisms. Pediatr Crit Care Med. 2003;4:465–470.

8. Nakamura KT, Sato Y, Erenberg A. Evaluation of a percutaneously placed 27-gauge central venous catheter in neonates weighing <1200 grams. J Parenter Enteral Nutr. 1990;14: 295–299.

9. Oloya RO, Feick HJ, Bozynski ME. Impact of venous catheters on packed red cells. Am J Perinatol. 1991;8:280–283.

10. Parshuram CS, Joffe AR. Prospective study of potassium-associated acute transfusion events in pediatric intensive care. Pediatr Crit Care Med. 2003;4:65–68.

11. Wilcox GJ, Barnes A, Modanlou H. Does transfusion using a syringe infusion pump and small gauge needle cause hemolysis. Transfusion. 1981;21:750–751.

12. Wong EC, Schreiber S, Criss VR, et al. Feasibility of red blood cell transfusion through small bore central venous catheters used in neonates. Pediatr Crit Care Med. 2004;5:69–74.

13. Gibson BE, Todd A, Roberts I, et al. Transfusion guidelines for neonates and older children. Br J Haematol. 2004;127:233–234.

14. Roseff SD, Luban NL, Manno CS. Guidelines for assessing appropriateness of pediatric transfusion. Transfusion. 2002;42: 1398–1413.

15. Shannon KM, Keith JF, Mentzer WC, et al. Recombinant human erythropoietin stimulates erythropoiesis and reduces erythrocyte transfusions in very low birth weight preterm infants. Pediatrics. 1995;95:1–8.
16. Bell EF, Strauss RG, Widness JA, et al. Randomized trial of liberal vs. restrictive guidelines for red blood cell transfusion in preterm infants. Pediatrics. 2005;115:1685–1691.

17. Luban NL, Strauss RG, Hume HA. Commentary on the safety of red cells preserved in extended-storage media for neonatal transfusion. Transfusion. 1991;31:229–235.

18. Strauss RG, Burmeister LF, Johnson K, et al. Feasibility and safety of AS-3 red blood cells for neonatal transfusions. J Pediatr. 2000;136:215–219.

19. Strauss RG. Controversies in the management of the anemia of prematurity using single-donor red blood cell transfusions and/or recombinant human erythropoietin. Transfus Med Rev. 2006;20:34–44.

20. Luban NL, Mikesell G, Sacher RA. Techniques for warming red blood cells packaged in different containers for neonatal use. Clin Pediatr (Phila). 1985;24:642–644.

21. Strauss RG, Bell EF, Snyder EL, et al. Effects of environmental warming on blood components dispensed in syringes for neonatal transfusions. J Pediatr. 1986;109:109–113.

22. Manno CS, Hedberg KW, Kim HC. Comparison of the hemostatic effects of fresh whole blood, stored whole blood, and components after open heart surgery in children. Blood. 1991;77: 930–936.

23. Mou SS, Giroir BP, Molitor-Kirsch EA, et al. Fresh whole blood versus reconstituted blood for pump priming in heart surgery in infants. N Engl J Med. 2004;351:1635–1644.

24. Brown KA, Bissonnette B, McIntyre B. Hyperkalaemia during rapid blood transfusion and hypovolaemic cardiac arrest in children. Can J Anaesth. 1990;37:747–754.

25. Martchenke J, Boshkov L. Heparin-induced thrombocytopenia in neonates. Neonatal Network. 2005;24:33–37.

26. Roberts IA, Murray NA. Neonatal thrombocytopenia. Curr Hematol Rep. 2006;5:55–63.

27. Moroff G, Friedman A, Robkin-Kline L, et al. Reduction of the volume of stored platelet concentrates for use in neonatal patients. Transfusion. 1984;2:144–146.

28. Haspel RL, Walsh L, Sloan SR. Platelet transfusion in an infant leading to formation of anti-D: implications for immunoprophylaxis. Transfusion. 2004;44:747–749.

29. Holman P, Blajchman MA, Heddle N. Noninfectious adverse effects of blood transfusion in the neonate. Trans Med Rev. 1995;9:277–287.

30. Sanchez R, Toy P. Transfusion related acute lung injury: a pediatric perspective. Pediatr Blood Cancer. 2005;45: 248–255.

31. Wu T-J, Teng R-J, Yau K-IT. Transfusion-related acute lung injury treated with surfactant in a neonate. Eur J Pediatr. 1996;155:589–591.

32. Kenton AB, Hegemier S, Smith EO, et al. Platelet transfusions in infants with necrotizing enterocolitis do not lower mortality but may increase morbidity. J Perinatol. 2005;25:173–177.

33. Engelfriet CP, Reesink HW, Klein HG, et al. International forum: granulocyte transfusions. Vox Sang. 2000;79:59–66.

34. Chalmers EA, Gibson BES. Clinical aspects of paediatric and perinatal transfusion: plasma products. Vox Sang. 1994;67: 54–58.

35. Boralessa H, Modi N, Cockburn H. RBC T activation and hemolysis in a neonatal intensive care population: implications for transfusion practice. Transfusion. 2002;42:1428–1434.

36. Ramasethu J, Luban NL. T activation. Br J Haematol. 2001;112:259–263.

37. Strauss RG, Burmeister LF, Johnson K, et al. Randomized trial assessing feasibility and safety of biologic parents as RBC donors for their preterm infants. Transfusion. 2000;40: 450–456.

38. Brune T, Garritsen H, Witteler R, et al. Autologous placental blood transfusion for the therapy of anemic neonates. Biol Neonate. 2002;81:236–243.

39. Eichler H, Schaible T, Richter E, et al. Cord blood as a source of autologous RBCs for transfusion to preterm infants. Transfusion. 2000;40:1111–1117.

40. Garritsen HS, Brune T, Louwen F. Autologous red cells derived from cord blood: collection, preparation, storage and quality controls with optimal additive storage medium (Sag-mannitol). Transfus Med. 2003;13:303–310.

41. Imura K, Kawahara H, Kitayama Y, et al. Usefulness of cord-blood harvesting for autologous transfusion in surgical newborns with antenatal diagnosis of congenital anomalies. J Pediatr Surg. 2001;36:851–854.

42. Dodd RY, Notari EP 4th, Stramer SL. Current prevalence and incidence of infectious disease markers and estimated window-period risk in the American Red Cross blood donor population. Transfusion. 2002;42:975–979.

43. Stramer SL, Glynn SA, Kleinman SH, et al. Detection of HIV-1 and HCV infections among antibodynegative blood donors by nucleic acid-amplification testing. N Engl J Med. 2004;351: 760–768.

44. Fang CT, Chambers LA, Kennedy J. Detection of bacterial contamination in apheresis platelet products: American Red Cross experience, 2004. Transfusion. 2005;45:1845–1852.

45. Mungai M, Tegtmeier G, Chamberland M, et al. Transfusion transmitted malaria in the United States from 1963 through 1999. N Engl J Med. 2001;344:1973–1978.

46. Ludlam CA, Turner ML. Managing the risk of transmission of variant Creutzfeldt Jakob disease by blood products. Br J Haematol. 2006;132:13-24.

42-Exchange Transfusions

Jayashree Ramasethu

The frequency with which postnatal exchange transfusions (ET) were performed decreased in the 1990s, resulting in significantly less experience among personnel performing the procedure (1, 2, 3). However, the re-emergence of kernicterus as a public health problem underscores the importance of ET as a treatment modality that could potentially prevent devastating neurodevelopmental complications (4, 5).

A. Definitions

• Exchange transfusion: Replacing an infant's blood with donor blood by repeatedly removing and replacing small aliquots of blood over a short time period

B. Indications

- Significant unconjugated hyperbilirubinemia in the newborn due to any cause, when intensive phototherapy fails or there is risk of kernicterus. Immediate ET is recommended if there are early signs of acute bilirubin encephalopathy (6). Figure 42.1 shows the total serum bilirubin levels at which ET is recommended for infants of 35 or more weeks' gestation.
- Indications for ET in more immature infants are variable and individualized (see Table 46.1). When performed for low bilirubin levels, ET is less effective than phototherapy in achieving prolonged reductions in total serum bilirubin in neonates with nonhemolytic jaundice (7).
- Alloimmune hemolytic disease of the newborn (HDN) (1)
 - For correction of severe anemia and hyperbilirubinemia
 - In addition, in infants with alloimmune HDN, ET replaces antibody-coated neonatal red cells with antigen-negative red cells that should have normal in vivo survival and also removes free maternal antibody in plasma.
- Severe anemia with congestive cardiac failure or hypervolemia (1, 8)
- Polycythemia

Although partial exchange transfusion with crystalloid or colloid reduces the packed cell volume and hyperviscosity in neonates with polycythemia, there appears to be no evidence of long-term benefit from the procedure (9).

- Disseminated intravascular coagulation (2)
- Congenital leukemia (10)
- Metabolic toxins
 - Hyperammonemia (11)
 - Organic acidemia (12)
 - Lead poisoning (13)
- Drug overdose or toxicity (14, 15)
- Removal of antibodies and abnormal proteins (16, 17)
- Neonatal sepsis or malaria (18, 19)

C. Contraindications

- When alternatives such as simple transfusion or phototherapy would be just as effective with less risk (7)
- When patient is unstable and the risk of the procedure outweighs the possible benefit

In infants with severe anemia, with cardiac failure or hypervolemia, partial ET may be useful to stabilize the patient's condition before a complete or double-volume ET is performed.

• When a contraindication to placement of necessary lines outweighs indication for ET. Alternative access should be sought if ET is imperative (20).

D. Equipment

- Infant care center (see Chapter 2)
 - Automatic and manually controlled heat source
 - Temperature monitor
 - Cardiorespiratory monitor
 - Pulse oximeter for oxygen saturation monitoring
- Resuscitation equipment and medication (immediately available)
- Infant restraints
- Orogastric tube
- Suctioning equipment
- Equipment for central and peripheral vascular access
- Blood warmer and appropriate coils (see Precautions)



 The dashed lines for the first 24 hours indicate uncertainty due to a wide range of clinical circumstances and a range of responses to phototherapy.

 Immediate exchange transfusion is recommended if infant shows signs of acute bilirubin encephalopathy (hypertonia, arching, retrocollis, opisthotonos, fever, high pitched cry) or if TSB is ≥5 mg/dL (85 µmol/L) above these lines.

- Risk factors isoimmune hemolytic disease, G6PD deficiency, asphyxia, significant lethargy, temperature instability, sepsis, acidosis.
- Measure serum albumin and calculate B/A ratio (See legend)
- Use total bilirubin. Do not subtract direct reacting or conjugated bilirubin

 If infant is well and 35-37 6/7 wk (median risk) can individualize TSB levels for exchange based on actual gestational age.

FIG. 42.1. Guidelines for exchange transfusion in infants 35 or more weeks gestation. (From American Academy of Pediatrics. Subcommittee on Hyperbilirubinemia. Clinical Practice Guideline. Management of hyperbilirubinemia in the newborn infant 35 or more weeks gestation. Pediatrics. 2004;114:297-316.)

- Sterile exchange transfusion equipment
 - Pre-assembled disposable set with special stopcock (Pharmaseal, Allegiance Healthcare Corporation, McGaw Park, IL, USA) or
 - o Nonassembled
 - 10- or 20-mL syringes Use a smaller syringe if aliquot per pass is smaller.
 - Two three-way stopcocks with locking connections
 - Waste receptacle (empty IV bottle or bag)
 - IV connecting tubing
- Appropriate blood product
- Syringes and tubes for pre- and postexchange blood tests

E. Precautions

- Stabilize infant before initiating exchange procedure.
- Do not start exchange procedure until personnel are available for monitoring and as backup for other emergencies.
- Use blood product appropriate to clinical indication. Use freshest blood available, preferably <5 to 7 days.
- Check potassium level of donor blood if patient has hyperkalemia or renal compromise.
- Monitor infant closely during and after procedure.
- Do not rush procedure.
 - May necessitate repeat ET if efficacy is decreased by haste.
 - Stop or slow procedure if patient becomes unstable.

- Use only thermostatically controlled blood-warming device that has passed quality control for temperature and alarms. Be sure to review operating and safety procedures for specific blood warmer. Do not overheat blood, i.e., beyond 38ŰC.
- Do not apply excessive suction if it becomes difficult to draw blood from line. Reposition line or replace syringes, stopcocks, and any adapters connected to line.
- Leave anticoagulated, banked blood in line or clear line with heparinized saline if the procedure is interrupted.
- Clear line with heparinized saline if administering calcium.
- F. Preparation for Total or Partial Exchange Transfusion
 - Blood product and volume Blood Product
 - Communicate with blood bank or transfusion medicine specialist to determine most appropriate blood product for transfusion.
 - Plasma-reduced whole blood or packed red cells reconstituted with plasma, with a packed cell volume adjusted to 0.5 to 0.60, is suitable for ET to correct anemia and hyperbilirubinemia (1, 21, 22 and 23).
 - Blood may be anticoagulated with citrate phosphate dextrose (CPD or CPDA1) or heparin. Additive anticoagulant solutions are generally avoided (21, 22, 23 and 24). If only RBCs stored in additive solutions are available, the additive solution may be removed by washing or by centrifugation and removal of the supernatant solution, prior to reconstitution of the red cells with plasma.
 - Blood should be as fresh as possible (<7 days).
 - Irradiated blood is recommended for all ET, to prevent graft-versus-host disease. There is a significant increase in potassium concentration in stored irradiated units, so irradiation should be performed as close to the transfusion as possible (<24 hours).
 - Standard blood-bank screening is particularly important, including sickle-cell preparation, HIV, hepatitis B, and cytomegalovirus (CMV) (see Chapter 41).
 - Donor blood should be screened for G-6-PD deficiency and HbS in populations endemic for these conditions (25).
 - In presence of alloimmunization, e.g., Rh, ABO, special attention to compatibility testing is necessary (1).
 - If delivery of an infant with severe HDN is anticipated, O Rh-negative blood crossmatched against the mother may be prepared before the baby is born.
 - Donor blood prepared after the infant's birth should be negative for the antigen responsible for the hemolytic disease, and should be cross-matched against the infant.
 - In ABO HDN, the blood must be type O and either Rh-negative or Rh-compatible with the mother and the infant. The blood should be washed free of plasma or have a low titer of anti-A or anti-B antibodies. Type O cells with AB plasma may be more effective, but this results in two donor exposures per ET (26).
 - In Rh HDN, the blood should be Rh-negative, and may be O group or the same group as the infant.
 - In infants with polycythemia, the optimal dilutional fluid for a partial ET is isotonic saline rather than plasma or albumin (27).

Volume of Donor Blood Required

- Whenever possible, use no more than equivalent of one whole unit of blood for each procedure, to decrease donor exposure.
- Quantity needed for total procedure = volume for the actual ET plus volume for tubing dead space and blood warmer (usually 25 to 30 mL)
- Double-volume ET for removal of bilirubin, antibodies, etc.:

- 2 infant's blood volume = 2 80 120 mL/kg
 - Infant's blood volume in preterm infant < 100 to 120 mL/kg, in term infant < 80 to 85 mL/kg
 - Exchanges approximately 85% of infant's blood volume (Fig. 42.2)
- Single volume ET: Exchanges approximately 60% of infant's blood volume (Fig. 42.2)
- Partial ET for correction of severe anemia:

Volume (mL)= $\underline{infant's blood volume \times (Hb desired-Hb initial)}$ Hb of PRBC-Hb initial

• Single-volume or partial ET for correction of polycythemia:

Volume (mL)= <u>infant's blood volume × desired HCT change</u> initial HCT

- Preparation of infant
 - Place infant on warmer with total accessibility and controlled environment. ET on small preterm infants may be performed in warm incubators, provided a sterile field can be maintained and lines are easily accessible.
 - Restrain infant suitably. Sedation and pain relief are not usually required. Conscious infants may suck on a pacifier during the procedure.



Fraction of Blood Volume Exchanged

FIG. 42.2. Graph depicting the effectiveness of exchange transfusion against the fraction of blood volume exchanged. The formula permits the calculation of the final hemoglobin.

- Connect physiologic monitors and establish baseline values (temperature, respiratory and heart rates, oxygenation)
- Empty infant's stomach.
 - Do not feed for 4 hours prior to procedure, if possible.

- Place orogastric tube and remove gastric contents; and leave on open drainage.
- Start IV line for glucose and medication infusion:
 - If exchange procedure interrupts previous essential infusion rate
 - If prolonged lack of oral or parenteral glucose will lead to hypoglycemia
 - Extra IV line may be necessary for emergency medications.
- Stabilize infant prior to starting exchange procedure; e.g., give packed-cell transfusion when severe hypovolemia and anemia are present, or modify ventilator or ambient oxygen when there is respiratory decompensation.
- Establish access for ET
 - Push pull technique: Central access usually through umbilical venous catheter.
 - Isovolumetric exchange with simultaneous infusion of donor blood through venous line and removal of baby's blood through arterial line. This technique may be better tolerated in sick or unstable neonates because there is less fluctuation of blood pressure and cerebral hemodynamics. The technique is also favored when only peripheral vascular access is available or preferred for various reasons (28, 29 and 30).
 - Infusion of donor blood may be through umbilical venous catheter or peripheral intravenous catheter.
 - Removal of baby's blood may be from umbilical arterial or venous catheter, or peripheral arterial catheter, usually a radial arterial line.
- Laboratory tests on infant's blood pre-exchange Tests are based on clinical indications.
 - Pre-exchange diagnostic studies. Note that diagnostic serologic tests on the infant, such as studies to evaluate unexplained hemolysis, antiviral antibody titers, metabolic screening, or genetic tests should be drawn prior to the ET.
 - Hemoglobin, hematocrit, platelets
 - Electrolytes, calcium, blood gas
 - Glucose
 - Bilirubin
 - Coagulation profile
- Prepare blood.
 - Verify identification of blood product (see Chapter 41).
 - Type and cross-match data
 - Expiration date
 - Donor and recipient identities
 - Attach blood administration set to blood-warmer tubing and to blood bag.
 - Allow blood to run through blood warmer.

G. Technique (See DVD for Animation of this Procedure)

Exchange Transfusion by Push Pull Technique through Special Stopcock with Preassembled Tray

- Read instructions provided by manufacturer carefully.
- Scrub as for major procedure. Wear mask, head cover, sterile gown, and gloves.
- Open preassembled equipment tray, using aseptic technique.
- Identify positions on special stopcock in clockwise rotation (Figs. 42.3 and 42.4). The direction that the handle is pointing indicates the port that is open to the syringe. The special stopcock allows clockwise rotation in the order used: (a) withdraw from patient; (b) clear to waste bag; (c) draw new blood; (d) inject into patient. Always rotate the handle in clockwise direction to follow the proper sequence, and keep connections tight.
 - Male adapter to umbilical or peripheral line
 - Female adapter to the extension tubing to which waste bag will be attached
 - Connect to blood tubing for attachment to blood-warmer coil.

• Neutral off position in which additives may be administered through rubber stopper (180 degrees from waste-receptacle port)



FIG. 42.3. Special four-way stopcock. A, male adapter to infant line; B, female adapter to waste container; C, attachment to blood tubing; D, off position (180 degrees from adapter to waste container), allowing injection through rubber-stoppered port below syringe. The stopcock is used in clockwise rotation when correctly assembled.

- Follow steps as provided by manufacturer to make all connections to blood and waste bags.
- With stopcock open to blood source, clear all air into syringe. Turn in clockwise direction 270 degrees and evacuate into waste.
- Turn stopcock to off and replace onto sterile field.
- Use pre-existing umbilical venous line or insert umbilical venous catheter, using sterile technique as described in Chapter 29 and attach spinal stopcock.
 - Consider CVP measurement, using pressure transducer, in unstable baby.
 - Place catheter in IVC and verify position by radiograph.
 - If catheter cannot be positioned in IVC, it may still be used cautiously in an emergency, when placed in the umbilical vein, if adequate blood return is obtained.



Fig. 42.4. A, B: ET using special four-way stopcock.

- Have an assistant document all vital signs, volumes, and other data on the exchange record.
- Check peripheral glucose levels every 30 to 60 minutes. Monitor cardiorespiratory status, continuous pulse oximetry. Determine blood gases as often as indicated by pre-existing clinical condition and stability.
- Draw blood for diagnostic studies
- Usual rate of removal and replacement of blood during the ET is aliquots of approximately 5 mL/kg over a 2- to 4-minute cycle.
- If infant is hypovolemic or has low CVP, start exchange with transfusion of aliquot (5 mL/kg) into catheter. If infant is hypervolemic or has high CVP, start by withdrawing precalculated aliquot.
- Remeasure CVP if indicated. Expect rise as plasma oncotic pressure increases, if CVP is low at start.
- Ensure that the stages of drawing and infusing blood from and into the infant are done slowly, taking at least a minute each to avoid fluctuations in blood pressure. Rapid fluctuations in arterial pressure in the push pull technique may be accompanied by changes in intracranial pressure (28, 31). Rapid withdrawal from the umbilical vein induces a negative pressure that may be transmitted to the mesenteric veins and contribute to the high incidence of ischemic bowel complications (32).
- Gently agitate the blood bag every 10 to 15 minutes to prevent red cell sedimentation, which may lead to exchange with relatively anemic blood towards the end of the exchange.
- Consider giving calcium supplement:
 - When hypocalcemia is documented
 - With symptoms or signs of hypocalcemia
 - Change in QTc interval
 - Agitation and tachycardia these symptoms are not reliably correlated with ionized calcium levels.

It is rarely necessary or advantageous to give calcium during an exchange if the infant is normocalcemic. When calcium is administered, the effect may last only a few minutes. Calcium will reverse the effect of the anticoagulant in the donor blood and may cause clotting of the line, so administration through a peripheral intravenous line is preferred. If calcium is given through an umbilical venous catheter, clear the line of donor blood prior to calcium administration with 0.9% NaCl. Give 1 mL of 10% calcium gluconate per kilogram body weight. Administer slowly, with careful observation of heart rate and rhythm. Clear line again with 0.9% NaCl.

• Perform calculated number of passes, until desired volume has been exchanged.

- Be sure there is adequate volume of donor blood remaining to infuse after last withdrawal, if a positive intravascular balance is desired.
- Clear umbilical line of banked blood and withdraw amount of infant's blood needed for laboratory testing, including re-cross-matching.
- Infuse IV fluids with 0.5 to 1 U heparin/mL of fluid through UVC if further ET are anticipated.
- \circ Total duration for double-volume ET: 90 to 120 minutes.
- Document procedure in patient's hospital record.

Exchange Transfusion Using a Single Umbilical Line and Two Three-Way Stopcocks in Tandem

The principles and techniques for using either the special stopcock or two three-way stopcocks in tandem are the same. It is important to ensure that all junctions are tight to produce a closed, sterile system. It is also essential to understand the working positions of the stopcocks before starting the exchange.

- Scrub as for major procedure. Wear sterile gown and gloves.
- Attach stopcock and tubing in sequence (Fig. 42.5).
 - Proximal stopcock
 - Umbilical catheter
 - IV extension tubing to sterile waste container
 - Distal stopcock
 - Tubing from blood-warming coil
 - 10- or 20-mL syringe
- Clear lines of air bubbles.
- Start exchange record.

Follow steps of push pull technique until exchange is completed.

Exchange Transfusion by Isovolumetric Technique (Central or Peripheral Line)

- Scrub as for major procedure.
- Select two sites for line placement, and insert as per Section 5, Vascular Access.
 - Venous for infusion
 - Umbilical venous catheter (preferred)
 - Peripheral IV cannula that is at least 23 gauge
 - Arterial for removal
 - Umbilical artery catheter (preferred)
 - Peripheral, usually radial if infant's size permits
- Connect arterial line to three-way stopcock.
 - Use short, connecting IV tubing to extend peripheral line.
 - Attach additional connecting tubing to stopcock and place into sterile waste container.



FIG. 42.5. Three-way stopcocks in tandem. Step 1: Stopcocks positioned for withdrawing blood from infant. Step 2: Stopcocks positioned for emptying withdrawn blood to waste container. Step 3: Stopcocks positioned for filling syringe from blood bag. Step 4: Stopcocks positioned for injecting blood into infant line.

• Attach empty 3- to 10-mL syringe to stopcock, for withdrawal of blood.

An additional stopcock may also be placed on this port so that a syringe of heparinized saline (5 U/mL) may be attached for use as needed. Be cautious about total volume infused.

- Connect venous line to single, three-way stopcock, which in turn connects to empty 5- to 10mL syringe and to blood-warming coil.
- Start exchange transfusion record.
- Withdraw and discard blood from arterial side at rate of 2 to 3 mL/kg/min, and infuse at same rate into venous side. Keep flow as steady as possible, and volumetrically equal for infusion and removal.
- Intermittently, flush arterial line with heparinized saline to clear.

The heparin solution remaining in tubing will be removed with next withdrawal, thus reducing significantly the total heparin dose actually received by the patient.

- Follow steps as for push–pull technique until exchange is complete.
- Total duration for isovolumetric ET: 45 to 60 minutes; may be longer in sick, unstable infants

H. Postexchange for All Techniques

- Continue to monitor vital signs closely for at least 4 to 6 hours.
- Rewrite orders; adjust any drug dosages as needed to compensate for removal by exchange (Table 42.1) (33, 34, 35).
- Keep infant NPO for at least 4 hours. Restart feeds cautiously if clinically stable. Monitor abdominal girth and bowel sounds every 3 to 4 hours for next 24 hours if exchange has been performed using umbilical vascular lines. Observe for signs of feeding intolerance.
- Monitor serum glucose levels every 2 to 4 hours for 24 hours.
- Repeat blood gases as often as clinically indicated.

	1	I el cent Loss		
Drug	One Volume	Two Volume		
Amikacin	7.1	13.8		
Ampicillin	7.7	14.7		
Carbamazepine	3.7	7.2		
Carbenicillin	5.6	10.9		
Colistin	18.7	33.9		
Diazepam	2.3	4.5		
Digoxina	1.2	2.4		
Furosemide	4.9	9.5		
Gentamicin	5.2	10.1		
Kanamycin	5.6	10.9		
Methicillin	10.1	19.1		
Oxacillin	19.6	35.4		
Penicillin G (crystalline)	6.0	11.6		
Penicillin G (procaine)	2.4	4.8		
Phenobarbital	6.4	12.3		
Phenytoin	3.1	6.2		
Theophylline ^a	17.8	32.4		
Tobramycin	10.3	19.6		
Vancomycin	5.7	11.0		

TABLE 42.1 Hypothetical Drug Loss by Exchange Transfusion Calculated by First-Order Elimination from a Single Compartment (1 â €" e^{-V/V}d) Percent Loss

V, plasma volume exchanged in liters; V_d , apparent volume of distribution. ^aWhole-blood volume used in calculation.

Source: From Lackner TE. Drug replacement following exchange transfusion. *J Pediatr.* 1982;100:813, with permission

- Measure serum ionized calcium levels and platelet counts in sick infants immediately after the ET and then as indicated.
- Repeat hemoglobin, hematocrit, and bilirubin measurements approximately 4 hours after exchange, and further as clinically indicated. A double-volume ET replaces 85% of the infant's blood volume, but eliminates only about 50% of the intravascular bilirubin. Equilibration of intra- and extravascular bilirubin, and continued breakdown of sensitized and newly formed red cells by persisting maternal antibody, results in a rebound of bilirubin levels following initial ET, and may necessitate repeated ET in severe HDN.

I. Complications

- Risk of death or permanent serious sequelae is estimated to be less than 1% in healthy infants, but as high as 12% in sick infants. There may be some uncertainty in ascribing adverse events to the ET procedure in infants who are already critically ill (36, 37, 38).
- Many of the adverse events are hematologic and biochemical laboratory abnormalities, which may be asymptomatic. The most common adverse effects noted during or soon after the ET, usually in infants who are preterm and/or sick include:
 - Apnea and/or bradycardia
 - Hypocalcemia
 - Thrombocytopenia (<50,000 in 10% of healthy infants, up to 67% of infants <32 weeks' gestational age)
 - Metabolic acidosis
 - Vascular spasm
- Complications reported from ET are related to the blood transfusion and to complications of vascular access (see Chapters 28, 29, 41).
- Potential complications include:
 - Metabolic: Hypocalcemia, hypo- or hyperglycemia, hyperkalemia
 - \circ Cardiorespiratory: Apnea, bradycardia, hypotension, hypertension
 - Hematologic: Thrombocytopenia, dilutional coagulopathy, neutropenia, disseminated intravascular coagulation
 - Vascular catheter-related: Vasospasm, thrombosis, embolization (see Chapter 28 and 29)
 - Gastrointestinal: Feeding intolerance, ischemic injury, necrotizing enterocolitis
 - o Infection: Omphalitis, septicemia

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References

1. Ramasethu J, Luban NLC. Alloimmune hemolytic disease of the newborn. In: Lichtman MA, Beutler E, Kipps TJ, et al. Williams' Hematology. 7th ed. New York: McGraw-Hill; 2006:751–766.

2. Funato M, Tamai H, Shimada S. Trends in neonatal exchange transfusions in Yodogawa Christian Hospital. Acta Paediatr Jpn. 1997;39:305–308.

3. Seidman DS, Paz I, Armon Y, et al. Effect of publication of the "Practice Parameter for the Management of Hyperbilirubinemia†on treatment of neonatal jaundice. Acta Paediatr. 2001;90:292–295.

4. Johnson LH, Bhutani VK, Brown AK. System based approach to management of neonatal jaundice and prevention of kernicterus. J Pediatr. 2002;140:396–403.

5. Ebbesen F. Recurrence of kernicterus in term and near term infants in Denmark. Acta Paediatr. 2000;89:1213–1217.

6. American Academy of Pediatrics. Subcommittee on Hyperbilirubinemia. Clinical Practice Guideline. Management of hyperbilirubinemia in the newborn infant 35 or more weeks gestation. Pediatrics. 2004;114:297–316.

7. Maisels MJ, Watchko JF. Treatment of jaundice in low birth weight infants. Arch Dis Child Fetal Neonatal Ed. 2003;88: 459–463.

8. Naulaers G, Barten S, Vanhole C, et al. Management of severe neonatal anemia due to fetomaternal transfusion. Am J Perinatol. 1999;16:193–196.

9. Dempsey EM, Barrington K. Short and long term outcomes following partial exchange transfusion in the polycythemic newborn. A systematic review. Arch Dis Child Fetal Neonatal Ed. 2006;91:F2–F6.

10. Fernandez MC, Weiss B, Atwater S, et al. Congenital leukemia: successful treatment of a newborn with t(5;11)(q31; q23). J Pediatr Hematol Oncol. 1999;21:152â \in "157.

11. Chen CY, Chen YC, Fang JT, et al. Continuous arteriovenous hemodiafiltration in the acute treatment of hyperammonaemia due to ornithine transcarbamylase deficiency. Ren Fail. 2000;22:823–836.

12. Aikoh H, Sasaki M, Sugai K, et al. Effective immunoglobulin therapy for brief tonic seizures in methylmalonic acidemia. Brain Dev. 1997;19:502–505.

13. Mycyk MB, Leikin JB. Combined exchange transfusion and chelation therapy for neonatal lead poisoning. Ann Pharmacother. 2004;38:821–824.

14. Sancak R, Kucukoduk S, Tasdemir HA, et al. Exchange transfusion treatment in a newborn with phenobarbital intoxication. Pediatr Emerg Care. 1999;15:268–270.

15. Osborn HH, Henry G, Wax P, et al. Theophylline toxicity in a premature neonateâ€"elimination kinetics of exchange transfusion. J Toxicol Clin Toxicol. 1993;31:639â€"644.

16. Pasternak JF, Hageman J, Adams MA, et al. Exchange transfusion in neonatal myasthenia. J Pediatr. 1981;99:644–646.

17. Dolfin T, Pomerance A, Korzets Z, et al. Acute renal failure in a neonate caused by the transplacental transfer of a nephrotoxic paraprotein: successful resolution by exchange transfusion. Am J Kidney Dis. 1999;34:1129–1131.

18. Gunes T, Koklu E, Buyukkayhan D, et al. Exchange transfusion or intravenous immunoglobulin therapy as an adjunct to antibiotics for neonatal sepsis in developing countries: a pilot study. Ann Trop Paediatr. 2006;26:39–42.

19. Virdi VS, Goraya JS, Khadwal A, Seth A. Neonatal transfusion malaria requiring exchange transfusion. Ann Trop Pediatr. 2003;23:205–207.

20. Burch M, Dyamenahalli U, Sullivan ID. Severe unconjugated hyperbilirubinemia with infradiaphragmatic total anomalous pulmonary venous connection. Arch Dis Child. 1993;68:608–609.
21. Goldenberg NA, Manco-Johnson MJ. Pediatric hemostasis and use of plasma components. Best Pract

Res Clin Haematol. 2006;19:143–155. 22. Murray NA, Roberts IAG. Neonatal transfusion practice. Arch Dis Child Fetal Neonatal Ed. 2004;89:101–107.

23. Petaja J, Johansson C, Andersson S, et al. Neonatal exchange transfusion with heparinised whole blood or citrated composite blood: a prospective study. Eur J Pediatr. 2000;159:552–553.

24. Win N, Amess P, Needs M, Hewitt PE. Use of red cells preserved in extended storage media for exchange transfusion in anti-k haemolytic disease of the newborn. Transfus Med. 2005; 15:157–160. 25. Kumar P, Sarkar S, Narang A. Acute intravascular haemolysis following exchange transfusion with G-6-PD deficient blood. Eur J Pediatr. 1994;153:98–99.

26. Yigit S, Gursoy T, Kanra T, et al. Whole blood versus red cells and plasma for exchange transfusion in ABO haemolytic disease. Transfus Med. 2005;15:313–318.

27. De Waal KA, Baerts W, Offringa M. Systematic review of the optimal fluid for dilutional exchange transfusion in neonatal polycythemia. Arch Dis Child Fetal Neonatal Ed. 2006;91: F7–F10.

28. Murakami Y, Yamashita Y, Nishimi T, et al. Changes in cerebral hemodynamics and oxygenation in unstable septic newborns during exchange transfusion. Kurume Med J. 1998;45:321–325.

29. Fok TF, So LY, Leung KW, et al. Use of peripheral vessels for exchange transfusions. Arch Dis Child. 1990;65:676–678.

30. Merchant RH, Sakhalkar VS, Rajadhyaksha SB. Exchange transfusion via peripheral vessels. Indian Pediatr. 1992;29: 457–460.

31. Van de Bor M, Benders MJ, Dorrepal CA, et al. Cerebral blood flow volume changes during exchange transfusions in infants born at or near term. J Pediatr. 1994;125:617–621.

32. Mintz AA, Vallbona C. A hazard of exchange transfusion in newborn infantsâ€"negative pressure in the umbilical vein. Pediatrics. 1960;26:661.

33. Ozkan H, Cevik N. Effect of exchange transfusion on elimination of antibiotics in premature infants. Turk J Pediatr. 1994;36:7–10.

34. Englund JA, Fletcher CV, Johnson D, et al. Effect of blood exchange on acyclovir clearance in an infant with neonatal herpes. J Pediatr. 1987;110:151.

35. Lackner TE. Drug replacement following exchange transfusion. J Pediatr. 1982;100:811.

36. Jackson JC. Adverse effects associated with exchange transfusion in healthy and ill newborns. Pediatrics. 1997;99:e7.

37. Keenan WJ, Novak KK, Sutherland JM, et al. Morbidity and mortality associated with exchange transfusion. Pediatrics. 1985;75(suppl):417–421.

38. Patra K, Storfer-Isser A, Siner B, et al. Adverse events associated with neonatal exchange transfusion in the 1990s. J Pediatr. 2004;144:626-631.

43-Removal of Extra Digits and Skin Tags

Leah Greenspan Hodor Khodayar Rais Bahrami

A. Indications

Removal of Nonfunctional Extra Digit

- Prevention of accidental avulsion or torsion around narrow base
- Cosmetic correction at parental request
- Prevention of traumatic amputation neuromas (1)

B. Contraindications

- Presence of illness in infant
- Presence of bleeding diathesis

This is an elective procedure that is painful when the clamp is applied. To prevent accidental avulsion of appendage if extra digit on a narrow base were to become entangled, apply a soft dressing or adhesive bandage until infant is stable enough for removal.

- Presence of any other hand anomaly when further surgical correction may be necessary (2)
- Base of extra digit >5 to 6 mm in width
- Bone crossing the isthmus between extra digit and hand

C. Equipment

- Iodophor antiseptic solution and swabs
- Straight mosquito hemostat
- Surgical silk suture, 3-0 or 4-0
- Fine or delicate scissors
- Adhesive bandage

D. Precautions

- Perform procedure only on stable, healthy infant.
 - Consider surgical evaluation for any questionable digit:
 - When base is >5 to 6 mm in width
 - When extra digit is on radial side of hand

- When clamping will not crush base to a thin, translucent layer indicative of hemostasis after excision
- When there appears to be a joint at the base
- When bony structures are present in the digit as confirmed by radiography (3)
- Apply hemostat to base of extra digit prior to placing ligature.
 - Allows closer amputation without residual bump
 - Allows faster autoamputation or removal of most of digit within a few hours

E. Technique

Removal of Nonfunctional Digit (Fig. 43.1)

- Cleanse digit and the surrounding skin with iodophor antiseptic. Allow to dry.
- Clamp hemostat as close to the base of extra digit as possible but without drawing up extra skin (Fig. 43.2A).
- Tightly tie suture around digit between hemostat and hand.
- Keep clamp in place until digit turns white for at least 5 minutes.
- Using as a cutting guide the edge of the hemostat farther from the hand, excise the digit (Fig. 43.2B).
- Remove hemostat and observe for hemostasis, leaving ligature in place. If there is any bleeding, reapply hemostat and ligature.
- Cover with an adhesive bandage until residual stump autoamputates.

Removal of Skin Tags (Fig. 43.3)

The removal of small skin tags follows essentially the same technique as for extra digits: Clamp close to base of lesion to achieve hemostasis, and apply ligature between hemostat and normal area. If the lesions are large or in critical areas, removal is best delayed beyond the neonatal period. Consider other diagnoses associated with skin tags (4).

F. Complications

- Hemorrhage
 - Failure to achieve complete hemostasis prior to excision
 - Loosening of ligature before blood supply is retracted
- Infection
- Inappropriate removal of digit in presence of related anomalies



FIG. 43.1. Nonfunctional extra digit on ulnar side of left hand.



FIG. 43.2. A: Place fine hemostat as close to base of extra digit as possible, and firmly secure ligature between clamp and hand. B: After finger turns white, excise digit tag outside hemostat, leaving ligature in place for autoamputation of residual stump.



FIG. 43.3. Skin tags of right ear and cheek. Removal of tags this large requires surgical excision rather than ligation for best result and may be associated with other malformations.

References

1. Leber GE. Surgical excision of pedunculated supernumerary digits prevents traumatic amputation neuromas. Pediatr Dermatol. 2003;20:108.

2. Nakamura J, Kanahara K, Endo Y, et al. Effective use of portions of the supernumerary digit to correct polydactyly of the thumb. Ann Plast Surg. 1985;15:7.

3. Gomella TL, Cunnigham MD, Eyal FG, Zenk KE. Newborn physical exam. In: Neonatology: Management, Procedures, On-Call Problems, Diseases, and Drugs. 4th ed. Stanford, CT: Appleton & Lange; 1999:35.

4. Jones KL. Perauricular tags or pits: frequent in. In: Jones KL, ed. Smith's Recognizable Patterns of Human Malformations. 6th ed. Philadelphia: Elsevier Saunders; 2006;899.

44-Circumcision

Mhairi G. MacDonald

A. Indications

Newborn male circumcision, one of the oldest formally recorded surgical procedures, remains controversial (1,2 and 3). Many physicians and lay people consider circumcision routine, but complications, although relatively rare, can be severe. Therefore, despite the perceived simplicity of the procedure, meticulous attention to anatomic landmarks, wound care, and follow-up is necessary.

A large French study compared the cost-effectiveness of circumcision and topical steroids as treatments for phimosis. The study concluded that topical pharmacologic treatment is effective in approximately 85% of patients, avoids the disadvantages, trauma, and potential complications of circumcision, and could reduce costs by 75% (4).

B. Contraindications

- Age <1 day (i.e., before complete physical adaptation to extrauterine life has occurred)
- Any current illness
- Prematurity (<37 weeks' gestation)
- Bleeding diathesis or family history of bleeding disorder
- Abnormality of urethra or penile shaft (foreskin may be essential for later reconstruction [e.g., hypospadias, chordee, very small penis])
- Local infection
- Lack of truly "informed parental consent (see Chapter 1)

C. Equipment

• Necessary for all methods

Sterile

- Gown and gloves
- Cup with antiseptic
- 4x4-in gauze pads
- Small, flexible, blunt probe
- Two straight mosquito hemostats

- Large, straight hemostat
- Tissue scissors

Nonsterile

- Materials for restraint
- Optional equipment
 - Local anesthetic: 1% lidocaine hydrochloride without epinephrine in a tuberculin syringe with a 1.2-cm x 27-gauge needle

Circumcision of neonates has frequently been used as a model to study the response of the newborn to pain (see Chapter 5). However, until recently, neonatal circumcision has been performed without anesthesia. Since the initial report by Kirya and Werthmann in 1978 (5), there have been reports of several controlled studies that have concluded that the use of dorsal penile nerve block is both effective and safe (5,6,7,8,9,10,11, and 12).

The effectiveness of EMLA (eutectic mixture of local anesthetics; lidocaine and prilocaine, Astra Pharma) 5% cream has also been studied (13,14 and 15). Conclusions from metaanalysis of data from refs. (12,13 and 14) led to the conclusion that EMLA cannot be recommended over other analgesic techniques with proven efficacy, such as regional nerve block with lidocaine. However, current data provide sufficient evidence to recommend routine use of EMLA for neonatal circumcision pain in settings in which no analgesics are routinely administered (15).

- Sterile fine-tipped marking pen
- Sterile gauze impregnated with petroleum jelly (e.g., Vaseline)
- Additional equipment for use with Gomco clamp

All equipment is sterile.

 Gomco circumcision clamp (Gomco Surgical Manufacturing Corp., Buffalo, NY, USA) (16), size 1 to 2 cm for average newborn glans (size range 1 to 3.5 cm)

Be sure to use a size that is large enough to protect the glans (17).

- o 11 scalpel blade and holder
- A small safety pin
- Additional equipment for use with Plastibell

All equipment is sterile.

Plastibell plastic cone (Hollister, Libertyville, IL, USA); available in presterilized packs; size range based on size of glans penis: 1.1, 1.3, and 1.5 cm. A linen suture is included in the pack (Fig. 44.1).

When selecting size, make sure that it is not so large that it allows proximal migration of the bell and excessive loss of penile skin nor so small that it could impair penile circulation.

• Scissors capable of cutting through plastic



FIG. 44.1. Plastibell with linen suture.

D. Precautions

- Obtain fully informed consent (see Chapter 1).
 - Explain expected course of circumcision to parents. When Plastibell is used, parents should be told to call their physician if ring has not fallen off within 10 days (18).
 - Be aware of laws pertaining to ritual circumcision (e.g., Jewish brit milah) (1,19,20) and the complications of the practice of orally suctioning the blood after cutting the foreskin (oral metzitzah) (21).
- Never circumcise at time of delivery. Circumcise long enough before discharge to allow adequate wound observation.
- Do not use local anesthetic containing epinephrine.
- Specifically locate coronal sulcus and urethral meatus.
- Make sure that inner epithelium is completely separated from glans penis and that prepuce can be retracted to visualize entire circumference of coronal sulcus.
- Never use electrocautery.
- Do not use circumferential dressing.
- Recheck wound prior to discharging patient and 1 to 2 weeks after circumcision. Residual skin should retract completely, and entire coronal sulcus must be visible to avoid postcircumcision adhesions, the most common complication.



FIG. 44.2. Penis is stabilized at angle of 20 to 25 degrees from midline. The formation of a lidocaine ring is shown (see text).

E. Technique

A complete description of formal surgical excision has been excluded from this edition because of the requirement to use sutures and the associated increased risk of bleeding compared with methods that involve crushing of tissue.

Ritual circumcisions are most commonly performed using a Mogen clamp. The method involves no dorsal incision or sutures (22); however, because the glans is not visible at the time of excision of the prepuce, there is potential for damage to the glans and urethra.

- Immobilize infant in supine position.
- Put on cap and mask.
- Scrub as for major procedure.
- Put on gown and gloves.
- Prepare skin with antiseptic, and drape.
- Perform penile dorsal nerve block if desired.
 - \circ Be familiar with anatomy of dorsal nerves of penis (Fig. 44.2) (5).

Although only the two dorsal penile nerves are targeted by the injection of lidocaine, the ventral penile nerve is also blocked by infiltration through the subcutaneous tissue. Some have advocated additional anesthesia ventrally, blocking the perineal nerves (a branch of the pudendal nerve) (23).

- Identify dorsal nerve roots at 10 o'clock and 2 o'clock positions.
- Identify by palpation the symphysis pubis and corpora cavernosa at the penile base.
- Estimate depth of pubic bone from penile base to indicate necessary depth of injection (should not exceed 0.5 cm).

Although the ideal area for infiltration corresponds to the 2 and 10 o'clock positions, 1 cm distal to the penile base, if the base is buried in pubic fat, the injection must be done at the junction of pubic and pelvic skin.

- Stabilize organ, with gentle traction, at angle of 20 to 25 degrees from midline.
- Pierce skin over one of dorsal nerves at penile root, and advance carefully posteromedially (0.25 to 0.5 cm) (Fig. 44.2) into subcutaneous tissue to avoid lodging in the erectile tissue.

After entering skin, needle should not meet resistance and tip should remain freely movable. If the tip of the needle is not freely mobile, it is probably embedded in the corpora cavernosum beneath the dorsal nerve and should be withdrawn slightly.

- Aspirate to rule out intravascular position.
- Slowly infiltrate area with 0.2 to 0.4 mL of lidocaine (never infiltrate as needle is advanced or withdrawn).

Arnett et al. (24) have reported good results using 0.2 mL of lidocaine.

• Repeat procedure at other dorsolateral position.

After infiltration, a small lidocaine ring forms (Fig. 44.2). The swelling is minimal and does not interfere with the circumcision procedure.

• Wait 3 to 5 minutes for analgesia.

Analgesia is usually obtained after 3 minutes and typically disappears within 20 to 30 minutes. However, there is individual variation, and testing of the prepuce with a hemostat is suggested prior to dissection.



FIG. 44.3. Circumcision. A: Marking the position of the coronal sulcus. B: Dilating the

preputial ring. C: Separating the prepuce from the glans penis. D: Grasping the prepuce with mosquito hemostats in preparation for the dorsal slit procedure. E: Dorsal slit.

- Locate coronal sulcus (Fig. 44.3A). Marking the position of the sulcus with ink on the skin of the penile shaft, prior to the procedure, is helpful in demarcating this vital landmark.
- Use mosquito hemostat to dilate preputial ring (Fig. 44.3B).
- Use blunt probe to separate inner epithelium of prepuce from glans penis (Fig. 44.3C).

Failure to do this completely may result in concealed penis (see G.3.c and G.13).

• Perform dorsal slit if desired.

This step is not mandatory as long as there is adequate separation of the glans from the prepuce.

- Grasp rim of prepuce on dorsal aspect with mosquito hemostats, approximately 2 to 4 mm apart (Fig. 44.3D).
- Visualize urethra.
- Place lower blade of large, straight hemostat between prepuce and glans to within 3 to 4 mm of corona, making sure to avoid urethra.
- Close hemostat for 5 to 10 seconds to crush foreskin in dorsal midline.
- Use tissue scissors to cut prepuce along crush line (Fig. 44.3E).
- Check that prepuce is freed from entire surface of glans. Complete separation if necessary.
- Complete circumcision using method of choice.
 - Use of circumcision clamp
 - Check clamp to ensure that all parts are present, fit well, and are in good working order.
 - Assemble clamp, ensuring that yolk (arm) articulates correctly with baseplate.
 - Draw prepuce backward gently to expose entire glans penis.
 - Break down all residual adhesions, and observe position of meatus. If meatus is abnormal, cease at this point.
 - Sponge glans dry with gauze swabs.
 - Select stud (bell) of adequate size (see C), and place over glans (Fig. 44.4A).
 - Pull prepuce over stud.
 - Approximate edge of dorsal slit. (A sterile safety pin may be used.)
 - Observe amount of skin remaining under baseplate for accuracy.

Proper placement of prepuce over stud is essential. Pulling too taut may lead to removal of excessive penile skin. Insufficient tension may lead to incomplete circumcision.




С

FIG. 44.4. Circumcision with a Gomco clamp. A: Placing the stud over the glans. B: Placing the baseplate of the clamp over the stud until the stud engages with the baseplate (inset). C: Gomco clamp in position for circumcision.

- Place baseplate of clamp over stud (with pin perpendicular to shaft of penis) so that prepuce is sandwiched between them (Fig. 44.4B).
- Continue to pull upward on stud until entire prepuce is drawn through baseplate and stud engages with baseplate.
- Hook yoke (arm) of clamp under side arms on shaft of stud and bolt firmly to baseplate, after checking position of prepuce between stud and baseplate (Fig. 44.4C).
- Remove safety pin.
- Wait 10 minutes.

Hemostasis is produced by pressure between baseplate and rim of stud. If the clamp is removed before 10 minutes has elapsed, wound edge hemostasis may be inadequate. If significant bleeding occurs during the procedure, remove the device and search for bleeding vesselâ€"avoid blindly placing sutures.

- Remove prepuce with scalpel held parallel to and flush with upper surface of baseplate. Never use electrocautery; however, use of an ultrasound dissection scalpel has been described as a safe alternative to electrocautery (25).
- Loosen bolt on clamp and remove.
- Optional: Dress with loose, noncircumferential sterile gauze impregnated with Vaseline.

Gough and Lawton (26) have shown that the addition of tincture of benzoin to the dressing adversely affected wound healing and the addition of topical antibiotic did not produce better results than those achieved with ordinary paraffin gauze.

- Apply tight diaper for 1 hour.
- For 24 hours after circumcision, check (or instruct parents to check) for bleeding, excessive swelling, and difficulty voiding.
- Until circumcised area is completely healed, do not immerse; give sponge bath.

Use of Plastibell

1.Follow Steps 3 to 5 of E.11.a.

2.Select bell of correct size (see C).

- Cone should fit snugly without pressure on glans
- Grooved rim of bell should be just distal to apex of dorsal slit.

3.If necessary, cut small segment out of cone so that it clears frenulum.

4.Hold prepuce firmly in place over cone (Fig. 44.5A).

5. Tie suture tightly around rim of bell so that prepuce is firmly compressed into groove.

6.Trim prepuce distal to ligature with tissue scissors. Use outer rim of cone as guide.

7.Break off cone handle. Tissue beneath ligature will atrophy and separate from bell in 5 to 8 days (maximum 10 to 12 days) (Fig. 44.5B).

8.Observe and care for circumcision as in Steps 17 and 18 of Gomco clamp (E.11.a).



FIG. 44.5. Circumcision with a Plastibell. A: The prepuce is pulled forward onto the bell. Inset: The prepuce is compressed into the groove by the circumferential suture. B: Appearance of the completed circumcision. F. Management of Postoperative Bleeding

Postoperative bleeding usually stems from inadequate hemostasis (e.g., unrecognized neonatal hepatitis [27] or hereditary clotting disorders). Rarely, anomalous vessels are responsible.

Continuous Ooze

• Apply manual pressure for 5 to 10 minutes.

Check that the string on the Plastibell is in place and is sufficiently tight.

- Assess bleeding site. If continued oozing:
 - Apply topical thrombin (Thrombostat) on absorbable gelatin sponge (Gelfoam) or oxidized cellulose (Oxycel, Surgicel); do not use circumferential dressing.
 - Silver nitrate and epinephrine have also been used topically to control bleeding. To avoid local ischemia or systemic effects, do not exceed a 1:100,000 concentration of epinephrine.

Active Hemorrhage or Uncontrolled Ooze

- Surgical assessmentâ€"ligation of bleeding vessel
- Consider underlying coagulopathy.

G. Complications (Fig. 44.6A-D)

The overall incidence of complications associated with circumcision ranges from approximately 0.2% to 7% (28,29 and 30).

- Hemorrhage (26,27,28,29,30, and 31)
- Infection (32,33, 34,35, 36, and 37)

More common with the Plastibell. Most are mild and respond to wet to dry dressings and Sitz baths, but fatalities have been reported (35).

- Local (5,32,33)
- Systemic (34,36)
- Necrotizing fasciitis (37)
- Incomplete circumcision (most common complication) (38,39 and 40)
 - Phimosis (40)
 - Skin bridge between penile shaft and glans (commonly due to inadequate skin removal and failure to visualize the corona on follow-up examination) (40,41)
 - \circ Concealed penis (see also G.13) (30,40,42,43 and 44)
- Trauma
 - Urethral laceration during dorsal slit procedure (avoided by keeping urethra in view at all times during the procedure)
 - Loss of penis (most commonly due to injuries related to cautery) (45,46)/amputation of glans (17)
 - Hypospadias/epispadias (47)
 - Cyanosis/necrosis of glans penis caused by overly tight Plastibell, misplaced sutures, or overtight circumferential bandage (30,40,48)
 - Urethrocutaneous fistula associated with use of Gomco clamp or Plastibell (most commonly caused by using a Plastibell or clamp of incorrect size or failure to recognize congenital megaloureter) (42,49,50)
- Urinary retention
 - Tight (or occlusive) dressing or glanular prolapse through ring of Plastibell (32,40,52,53 and 54)
 - Meatal stenosis resulting from urethral meatitis (5,40)
- Inflammation/ulceration of meatus (1,5,54,55,56,57, and 58)
- Circumcision of hypospadias (40,52)
- Chordee (40); most commonly is the result of dense ventral scarring from inflammation; may be due to removal of excess skin from shaft or secondary to a skin bridge
- Inclusion cyst of prepuce (52)
- Lymphedema (40,52,59)
- Venous stasis (60)
- Displacement with lodging of Plastibell around penile shaft or glans penis (18,61,62 and 63)
- Death
 - Anesthetic (1,63,64)
 - Infection (65)
- Wound separation/removal of excess skin (Fig. 44.6) (30,52,61,66)

Buried penis is usually the result of inappropriate circumcision in a chubby baby with a small or concealed penis. Excessive removal of skin should be treated with application of antiseptic (iodophor) daily and not with grafting or burying the penis in scrotum. The skin will grow back.

- Recurrence of pneumothorax (67)
- Reaction to epinephrine used to control bleeding (68)
 - Tachycardia
 - Local vasospasm (may lead to necrosis of the glans)
- Complications due to local anesthetic
 - Methemoglobinemia has been reported following exposure to prilocaine, procaine, benzocaine, and lidocaine (69).
 - Hematoma (70); those reported in neonates have resolved spontaneously.
 - Seizures (71)
- Mechanical problems with Gomco clamp (72)
 - $\circ \quad Loss \ of \ a \ part$
 - Warping of the plate after multiple use
 - Breakage of arm during tightening
 - Grooves and nicks in bell at junction of bell and plate



FIG. 44.6. Complication of circumcision. A: Glans injury 6 months after circumcision. B: Trapped penis following contraction of wound after circumcision. C: Penile amputation following cautery injury during circumcision. D: Cicatrix following circumcision.

References

1. Gairdner D. The fate of the foreskinâ€"a study of circumcision. Br Med J. 1949;2:1433.

2. Poland RL. The question of routine circumcision. N Engl J Med. 1990;322:1312.

3. Hutcheson JC. Male neonatal circumcision: indications, controversies and complications. Urol Clin North Am. 2004;31:461, viii.

4. Berdeu D, Sauze L, Ha-Vinh P, et al. Cost-effectiveness analysis of treatments for phimosis: a

comparison of surgical and medical approaches and their economic effect. Br J Urol Int. 2001;87:239.

5. Kirya C, Werthmann MW Jr. Neonatal circumcision and penile dorsal nerve block—a painless procedure. J Pediatr. 1978;92:998.

6. Pelosi AP, Apuzzio J. Making circumcision a painless event. Contemp Pediatr. 1985;Jan:85.

7. Maxwell LG, Yaster M, Wetzel RC, et al. Penile nerve block for newborn circumcision. Obstet Gynecol. 1987;70:415.

8. Sfez M, Lemapihan Y, Mazoit X, et al. Local anesthetic concentrations after penile nerve block in children. Anesth Analg. 1990;71:423.

9. Soh CR, Ng SB, Lim SL. Dorsal penile nerve block. Pediatr Anaesth. 2003;13:329.

 Lander J, Brady-Fryer B, Nazarali S, Muttitt S. Three interventions for circumcision pain. International Association for the Study of Pain. 8th World Congress on Pain, Vancouver, Canada; 1996:186[abstr 256].
 Garry DJ, Swoboda E, Elimian A, Figueroa R. A video study of pain relief during newborn male circumcision. J Perinatol. 2006;26:106.

12. Taddio A, Stevens B, Craig K, et al. Efficacy and safety of lidocaine–prilocaine cream for pain during circumcision. N Engl J Med. 1997;336:1197.

13. Benini F, Johnston CC, Faucher D, Aranda JV. Topical anesthesia during circumcision in newborn infants. JAMA. 1993;270:850.

14. Brady-Fryer B, Wiebe N, Lander JA. Pain relief for neonatal circumcision. Cochrane Database Syst Rev. 2004;18:CD004217.

15. Taddio A, Ohlsson A (Reviewers). Lidocaine–prilocaine cream (EMLA) to reduce pain in male neonates undergoing circumcision. www.nichd.nih.gov/cochrane/Taddio/Taddio1.htm; e-mail: anna.taddio@uteronto.ca.

16. Yellen HS. Bloodless circumcision of the newborn. Am J Obstet Gynecol. 1935;30:146.

17. Essid A, Hamazaoui M, Sahli S, Houissa T. Glans reimplantation after circumcision accident. Prog Urol. 2005;15:745.

18. Rubenstein MM, Basen WM. Complication of circumcision done with a plastic bell clamp. Am J Dis Child. 1968; 116:381.

19. Gottilieb N. A Jewish Child Is Born. New York: Block; 1960.

20. Glass JM. Religious circumcision: a Jewish view. Br J Urol Int. 1999;83(suppl 1):17.

21. Gesundheit B, Grisaru-Soen G, Greenberg D, et al. Neonatal genital herpes simplex type 1 infection after Jewish ritual circumcision: modern medicine and religious tradition. Pediatrics. 2004;114:e259.

22. Dubrisin R, Zaprudsky P. Circumcising neonates with the Mogen clamp. Contemp OB/Gyn. 1991;36:79.

23. Serour F, Mori J, Barr J. Optimal regional anesthesia for circumcision. Anesth Analg. 1994;79:129.

24. Arnett RM, Jones JS, Horger EO. Effectiveness of 1% lidocaine dorsal penile nerve block in infant circumcision. Am J Obstet Gynecol. 1990;163:1074.

25. Fette A, Schleef J, Haberlik A, Seebacher U. Circumcision in pediatric surgery using an ultrasound dissection scalpel. Technol Health Care. 2000;8:75.

26. Gough DCS, Lawton N. Circumcisionâ€" which dressing? Br J Urol. 1990;65:418.

27. Hiss J, Horowitz A, Kahana T. Fatal haemorrhage following male ritual circumcision. J Clin Forensic Med. 2000;7:32.

28. Wiswell TE, Geschke DW. Risks from circumcision during the first month of life compared with those for uncircumcised boys. Pediatrics. 1989;83:1011.

29. Moreno CA, Realini JP. Infant circumcision in an outpatient setting. Texas Med. 1989;85:37.

30. Gee WF, Ansell JS. Neonatal circumcision: a ten year overview with comparison of the Gomco clamp and the Plastibell device. Pediatrics. 1976;58:824.

31. Patel H. The problem of routine circumcision. Can Med Assoc J. 1966;95:576.

32. Frand M, Berant N, Brand N, et al. Complications of ritual circumcision in Israel. Pediatrics. 1974;54:521.

33. Rosenstein JL. Wound diphtheria in the newborn infant following circumcision. J Pediatr. 1941;18:657.

34. Rubin LG, Lanzkowsky P. Cutaneous neonatal herpes simplex infection associated with ritual circumcision. Pediatr Infect Dis J. 2000;19:266.

35. Gearhart JP, Callan NA. Complications of circumcision. Contemp Ob/Gyn. 1986; 27: 57.

- 36. Kirkpatrick BV, Eitzman DV. Neonatal septicemia after circumcision. Clin Pediatr. 1974;13:767.
- 37. Woodside JR. Necrotizing fasciitis after neonatal circumcision. Am J Dis Child. 1980;134:301.
- 38. Leitch IOW. Circumcisionâ€"a continuing enigma. Aust Pediatr J. 1970;6:59.

39. Wynder EL, Licklider SD. The question of circumcision. Cancer. 1960;13:442.

40. Kaplan GW. Circumcisionâ€"an overview. Curr Probl Pediatr. 1977;7:3.

41. Klauber GT. Circumcision and phallic fallacies, or the case against routine circumcision. Conn Med. 1973;37:445.

42. Byars LT, Triers WC. Some complications of circumcision and their surgical repair. Arch Surg. 1958;76:477.

43. Talarico RD, Jasaitis JE. Concealed penis: a complication of neonatal circumcision. J Urol. 1973;110:732.

44. Trier WC, Drach GW. Concealed penisâ€"another complication of circumcision. Am J Dis Child. 1973;125:276.

45. Izzidien AY. Successful reimplantation of a traumatically amputated penis in a neonate. J Pediatr Surg. 1981;16:202.

- 46. Money J. Ablatio penis: normal male infant sex-reassignment as a girl. Arch Sex Behav. 1975;4:65.
- 47. McGowan AJ Jr. Letter to the editor: a complication of circumcision. JAMA. 1969;207:2104.

48. Rosefsky JB. Glans necrosis as a complication of circumcision. Pediatrics. 1967;39:774.

49. Limaye RD, Hancock RA. Penile urethral fistula as a complication of circumcision. J Pediatr. 1968;72:105.

50. Shiraki IW. Congenital megalourethra with urethrocutaneous fistula following circumcision: a case report. J Urol. 1973; 109:723.

51. Cook A, Koury AE, Bagli DJ, et al. Use of buccal mucosa to simulate the coronal sulcus after traumatic penile amputation. Urology. 2005;66:1109.

52. Shulman J, Ben-Hur N, Neuman Z. Surgical complications of circumcision. Am J Dis Child. 1965;107:149.

53. Berman W. Urinary retention due to ritual circumcision. Pediatrics. 1975;56:621.

54. Horowitz J, Sussheim A, Scalettar HE. Abdominal distention following ritual circumcision. Pediatrics. 1976;57:579.

55. Pearce I. Retention of urine: an unusual complication of the Plastibell device. Br J Urol Int. 2000;85:560. 56. Aley MC. Circumcision. JAMA. 1970;214:2195.

57. Gallagher AGP. Complications of circumcision. Br J Urol. 1972;44:720.

58. Mackenzie AR. Meatal ulceration following neonatal circumcision. Obstet Gynecol. 1966;28:221.

59. Yildirim S, Taylan G, Akoz T. Circumcision as an unusual cause of penile lymphedema. Ann Plast Surg. 2003;50:665.

60. Ly L, Sankaran K. Acute venous stasis and swelling of the lower abdomen and extremities in an infant after circumcision. Can Med Assoc J. 2003;169:216.

61. Johnsonbaugh RE, Meyer BP, Catalano JD. Complication of a circumcision performed by a plastic bell clamp. Am J Dis Child. 1969;118:781.

62. Malo T, Bonforte RJ. Hazards of plastic bell circumcision. Obstet Gynecol. 1969;33:869.

63. Wright JE. Non-therapeutic circumcision. Med J Aust. 1967;1: 1083.

64. Mal Herbe WOF. Injuries to the skin of the male genitalia in South Africa. S Afr Med J. 1975;49:147.

65. Grossman E, Posner NA. Surgical circumcision of neonates: a history of its development. Obstet Gynecol. 1981;58:241.

66. Van Duyn J, Warr WS. Excessive penile skin loss from circumcision. J Med Assoc Ga. 1962;51:394. 67. Auerbach MR, Scanlon JW. Recurrence of pneumothorax as a possible complication of elective circumcision. Am J Obstet Gynecol. 1978;132:583.

68. Denton J, Schriener RL, Pearson J. Circumcision complication: reaction to treatment of local hemorrhage with topical epinephrine in high concentration. Clin Pediatr. 1978;17: 285.

69. Couper RT. Methaemoglobinaemia secondary to topical lignocaine/prilocaine in a circumcised neonate. J Paediatr Child Health. 2000;36:406.

70. Fontaine P, Toffler WL. Dorsal nerve block for newborn circumcision. Am Fam Practitioner. 1991;43:1327.

71. Moran LR, Hossain T, Insoft RM. Neonatal seizures following lidocaine administration for elective circumcision. J Perinatol. 2004;24:395.

72. Feinberg AN, Blazek MA. Mechanical complications of circumcision with a Gomco clamp. Am J Dis Child. 1988;142: 813.

45-Drainage of Superficial Abscesses

An N. Massaro Khodayar Rais-Bahrami

A. Definitions

A superficial abscess is

- A localized collection of pus that causes fluctuant soft tissue swelling and may have associated erythema and induration (Fig. 45.1) (1, 2,3 and 4)
- Usually caused by invasion of local bacterial flora (1) or direct inoculation, i.e., animal bites ((5)) or intravenous access/skin piercing (6,7 and 8)
- A result of bacterial organisms that cause necrosis, liquefaction, accumulation of leukocytes and debris, followed by loculation and walling off of pus (9)
- **B.** Indications
 - To establish free drainage of contents from a superficial abscess

Surgical incision and drainage is the definitive treatment for soft tissue abscesses. Antibiotic therapy alone is ineffective in the setting of localized abscess and may even be unnecessary as an adjuvant to complete surgical drainage (1,2,10,11).

- To identify pathogens and direct antimicrobial therapy if needed (12, 13,14 and 15)
- To differentiate infectious from noninfectious lesions (13,16,17)

C. Contraindications

- Carefully identify and avoid:
 - o Cephalohematoma
 - Hemangioma
 - Cystic hygroma
 - Encephalocele

- Avoid premature incision and drainage of abscesses that have not yet fully matured, i.e., in the initial stages of induration and inflammation prior to formation of pus (9). This may lead to:
 - \circ A noncurative intervention
 - Possible extension of infectious process
 - o Bacteremia

This may be avoided by the use of ultrasound with or without diagnostic needle aspiration (18,19). D. Equipment

Sterile

- Gloves and gown
- Antiseptic swabs or cup containing antiseptic solution
- 1-mL syringe
- Nonbacteriostatic, isotonic saline without preservative
- 23-gauge needle
- 2 x 2-in gauze squares
- Scalpel with no. 11 blade
- Cotton-tipped culture swab
- Mosquito hemostat
- 2-in, fine-mesh, plain gauze

Nonsterile

- Ethyl chloride spray as topical anesthetic (For larger lesions, local anesthesia with lidocaine may be used.)
- Mask and cap
- Adhesive tape

E. Precautions

- Use appropriate isolation techniques to safeguard other infants.
- Obtain blood cultures after drainage.
- Do not suture abscess cavity following incision and drainage.
- Débride all tissue undergoing putrefaction and digestion thoroughly (4).
- Make skin incisions:
 - \circ $\,$ Conform with skin creases/natural folds to minimize scar formation
 - $_{\odot}$ Large enough to allow for proper débridement and drainage
 - Simple linear–cruciate or elliptical skin incisions may result in more unsightly scar formation ((9)).
- For abscesses in cosmetic areas, areas under significant skin tension (i.e., extensor surfaces), or areas with extensive scar tissue (i.e., sites of prior drainage procedures), a stab incision or needle aspiration alone may be preferable. (This may require multiple decompressions and/or delayed complete incision and drainage if reaccumulation occurs.)
- Care should be taken in areas with abundant vascular and neural structures, such as the groin, posterior knee, antecubital fossa, and neck ((3)).
- If foreign body is suspected, a radiograph should be obtained ((9)).



FIG. 45.1. Superficial abscess in the site of a Broviac central venous line insertion in the left anterior chest wall.

- F. Technique (1,4,7,8)
 - Spray roof of abscess with ethyl chloride until skin becomes white. (If local anesthesia is required, lidocaine can be injected subcutaneously with a 25-gauge needle into the dome of the abscess).
 - Prepare as for major procedure if abscess is to be drained, or for minor procedure if needle aspiration alone is to be performed (see Chapter 4).
 - Prepare local area with antiseptic (e.g., iodophor).
 - Aspiration [may be performed in combination with incision and drainage for confirmation of presence of pus and collection of material for culture, or alone if abscess is in area where incision is not preferable (see E.6)].
 - Attach sterile needle to syringe.
 - Insert needle into pustule, abscess cavity, or advancing border of cellulitis.
 - Aspirate the material deep within the lesion.
 - If no material is aspirated, inject 0.1 to 0.2 mL of nonbacteriostatic saline and withdraw immediately.
 - Process aspirated material immediately: Gram stain and culture for anaerobic and aerobic organisms; Giemsa stain for suspected herpes. Perform other special stains as warranted.
 - Incision and drainage
 - Insert scalpel blade and incise at point of maximum fluctuance. The size of the incision should be as small as possible yet allow for continued adequate drainage (i.e., length of the abscess cavity).
 - Obtain specimen for culture with cotton-tipped applicator, if not obtained by prior aspiration with syringe and needle.
 - Evacuate exudate from abscess with gentle pressure from finger or hemostat wrapped in gauze. Use caution when probing abscess with finger in cases of suspected retained foreign bodies or fragmentsâ€"for this reason, hemostat wrapped in gauze is the preferred method ((9)).

- If necessary, insert mosquito hemostat into abscess cavity and spread blades to break septa and to release remaining collections of pus (Fig. 45.2A). Recognize that this may cause discomfort and should be done rapidly.
- Lavage area with sterile saline to remove residual pus (optional).
- If indicated, insert plain, ¹/₂-in gauze into abscess cavity to stop bleeding and/or to serve as a wick to promote drainage (Fig. 45.2B).
- Apply dry, sterile dressing.
- Remove half of gauze packing in 24 hours and remainder within 48 hours. (Some larger wounds may require multiple packing changes).
- Check abscess wound, and apply sterile warm soaks for 20 to 30 minutes, three times a day, until healing has commenced, as indicated by:
 - Cessation of drainage
 - Formation of granulation tissue
 - Resolution of local tissue inflammation

G. Complications

- Introduction of infection into sterile abscess or hematoma
- Local bleeding
- Injury to blood vessels, nerves, or tendons (deep to abscess cavity) (3)
- Incomplete drainage with recurrent abscess formation (2,4)
- Systemic infection (20,21)
- Scar formation at drainage site, requiring skin graft (22)
- Reduction of breast size following incomplete drainage of breast abscess (23)



FIG. 45.2. Drainage of a superficial abscess. A: Breaking the septa with a clamp. B: Packing the wound.

References

1. Meislin HW, McGehee MD. Management and microbiology of cutaneous abscesses. JACEP. 1978;7:186.

2. Meislin HN, Lerner SA, Graves MH, et al. Cutaneous abscesses: anaerobic and aerobic bacteriology and outpatient management. Ann Intern Med. 1977;87:145.

3. Albom M. Surgical gems: surgical management of a superficial cutaneous abscess. J Dermatol Surg Oncol. 1976;2:120.

4. MacFie J, Harvey J. The treatment of acute superficial abscesses: a prospective clinical trial. Br J Surg. 1977;64:264.

5. Brook I. Microbiology and management of human and animal bite wound infections. Primary Care. 2003;30:25.

6. Schnall SB, Holton PD, Lilley JC. Abscesses secondary to parenteral abuse of drugs. J Bone Joint Surg Am. 1994;76:1526.

7. Murphy EL, DeVita D, Liu H, et al. Risk factors for skin and soft tissue abscess among injection drug users: a case-control study. Clin Infect Dis. 2001;33:35.

8. Folz BJ, Lippert BM, Kuelkens C, Werner JA. Hazards of piercing and facial body art: report of three patients and literature review. Ann Plast Surg. 2000;45:374.

9. Butler KH. Incision and drainage. In: Roberts JR, Hedges JR, eds. Clinical Procedures in Emergency Medicine. 4th ed. Philadelphia: WB Saunders; 2004:717.

10. Llera JL, Rios AM, Aten MF, et al. Treatment of cutaneous abscess: a double-blind clinical study. Ann Emerg Med. 1985; 14:15.

11. Lee MC, Rios AM, Aten MF, et al. Management and outcome of children with skin and soft tissue abscesses caused by community-acquired methicillin-resistant Staphylococcus aureus. Pediatr Infect Dis J. 2004;23:123.

12. Goetz J, Tafari N, Boxerbaum B. Needle aspiration in Haemophilus influenzae type B cellulitis. Pediatrics. 1974;54: 504.

13. Rudoy R, Nakashima G. Diagnostic value of needle aspiration in Haemophilus influenzae type B cellulitis. J Pediatr. 1979;94:924.

14. Allen CH, Patel B, Endom EE. Primary bacterial infections of the skin and soft tissues: changes in epidemiology and management. Clin Ped Emerg Med. 2004;5:246.

15. Zetola N, Francis JS, Nuermberger EL, Bishai WR. Community-acquired methicillin-resistant Staphylococcus aureus: an emerging threat. Lancet Infect Dis. 2005;5:275.

16. Jarratt M, Ramsdell W. Infantile acropustulosis. Arch Dermatol. 1979;115:834.

17. Kahn G, Rywlin A. Acropustulosis of infancy. Arch Dermatol. 1979;115:831.

18. Cardinal E. Bureau NJ, Aubin B, Chhem RK. Role of ultrasound in musculoskeletal infections. Radiol Clin North Am. 2001;39:191.

19. Loyer EM, DuBrow RA, David CL, et al. Imaging of superficial soft-tissue infections: sonographic findings in cases of cellulites and abscess. AJR: Am J Roentgenol. 1996;166:149.

20. Fine BC, Sheckman PR, Bartlett JC. Incision and drainage of soft-tissue abscesses and bacteremia [letter]. Ann Intern Med. 1985;103:645.

21. Blick PWH, Flowers MW, Marsden AK, et al. Antibiotics in surgical treatment of acute abscesses. Br Med J. 1980;281:111.

22. Feder H, McLean WC, Moxon R. Scalp abscess secondary to fetal scalp electrode. J Pediatr. 1976;89:808.

23. Rudoy R, Nelson J. Breast abscess during the neonatal period. Am J Dis Child. 1975;129:1031.

46-Phototherapy

Sepideh Nassabeh-Montazami

Phototherapy is the most common therapeutic intervention used for the treatment of hyperbilirubinemia (1). The process of phototherapy causes three reactions: photo-oxidation, configurational and structural isomerization of the bilirubin molecule, leading to polar, water-soluble photoproducts that can be excreted in bile and urine without the need for conjugation or further metabolism (2).

The aim of phototherapy is to reduce serum bilirubin levels in order to decrease the risk of acute bilirubin encephalopathy and the more chronic side effect of bilirubin toxicity, kernicterus (3).

A. Indications

- Clinically significant indirect hyperbilirubinemia. Indications to start phototherapy in babies with hyperbilirubinemia can vary depending on gestational age, birthweight, hours of life, presence of hemolysis, and other risk factors such as acidosis and sepsis (3,4).
- The total serum bilirubin (TSB) level must be considered when making the decision to commence treatment, as there is significant variability in laboratory measurement of direct bilirubin levels.
- The American Academy of Pediatrics has published clinical practice guidelines for phototherapy in newborn infants at 35 weeks' or more gestation (3) (Fig. 46.1).
- These guidelines do not apply to preterm infants less than 35 weeks' gestation. Preterm infants are at higher risk of developing hyperbilirubinemia compared to term infants. The decision to initiate phototherapy in this group of infants remains variable and highly individualized. The management of hyperbilirubinemia in extremely preterm and low-birthweight infants has been recently reviewed (4,5) (Table 46.1).
- Prophylactic phototherapy, often used for preterm infants with skin bruising, is ineffective until bilirubin appears in the skin.

B. Contraindications

- Congenital porphyria or a family history of porphyria is an absolute contraindication to the use of phototherapy. Severe purpuric bullous eruptions have been described in neonates with congenital erythropoietic porphyria treated with phototherapy (6).
- Concomitant use of drugs or agents that are photosensitizers is also an absolute contraindication (7).
- Concurrent therapy with metalloporphyrin heme oxygenase inhibitors has been reported to result in mild transient erythema (8).
- Although infants with cholestatic jaundice may develop the "bronze baby syndrome†when exposed to phototherapy (see Complications), the presence of direct hyperbilirubinemia is no longer considered to be a contraindication (3). However, because the products of phototherapy are excreted in the bile, the presence of cholestasis may decrease the effectiveness of phototherapy.

C. Equipment

In order to have an understanding of the equipment available for phototherapy, it is necessary to be familiar with the terminology involved.

- Spectral qualities of the delivered light (wavelength range and peak). Bilirubin absorbs visible light within the wavelength range of 400 to 500 nm, with peak absorption at 460 ± 10 nm considered to be the most effective (2).
- Irradiance (intensity of light), is expressed as watts per square centimeter (W/cm²). This refers to the number of photons received per square centimeter of exposed body surface area.

• Spectral irradiance is irradiance that is quantitated within the effective wavelength range for efficacy and is expressed as $\hat{A}\mu W/cm^2/nm$ (9). This is measured by various commercially available radiometers. Specific radiometers are generally recommended for each phototherapy system, because measurements of irradiance may vary depending on the radiometer and the light source (3).

A variety of phototherapy equipment devices exist and may be free-standing, attached to a radiant warmer, wall-mounted, suspended from the ceiling, or fiber-optic systems. These in turn may contain various light sources to deliver the phototherapy, which may be categorized as:



Use total bilirubin. Do not subtract direct reacting or conjugated bilirubin.

 Risk factors = isoimmune hemolytic disease, G6PD deficiency, asphyxia, significant lethargy, temperature instability, sepsis, acidosis, or albumin < 3.0g/dL (if measured)

For well infants 35-37 6/7 wk can adjust TSB levels for intervention around the medium risk line. It is an option to
intervene at lower TSB levels for infants closer to 35 wks and at higher TSB levels for those closer to 37 6/7 wk.

It is an option to provide conventional phototherapy in hospital or at home at TSB levels 2-3 mg/dL (35-50mmol/L)

below those shown but home phototherapy should not be used in any infant with risk factors.

FIG. 46.1. Guidelines for phototherapy in hospitalized infants of 35 or more weeks' gestation. (From American Academy of Pediatrics Subcommittee on Hyperbilirubinemia. Management of hyperbilirubinemia in the newborn infant 35 or more weeks of gestation. Pediatrics. 2004;114:297-316.)

- Fluorescent tubes
- Halogen bulbs
- Fiber-optic light combinations used in pads, blankets, or spotlights
- o High-intensity light-emitting diodes

The clinician is therefore faced with a vast array of equipment to choose from and must be aware of advantages and disadvantages of each type.

	Total Bilirubin Level (mg/dL/µmol/L) Exchange Transfusion			
Gestational Age (wk)				
	Phototherapy	Sick ^a	Well	
36	14.6 (250)	17.5 (300)	20.5 (350)	
32	8.8 (150)	14.6 (250)	17.5 (300)	
28	5.8 (100)	11.7 (200)	14.6 (250)	
24	4.7 (80)	8.8 (150)	11.7 (200)	

TABLE 46.1 Guidelines for Use of Phototherapy and Exchange Transfusion in Preterm InfantsBased on Gestational Age

^aRhesus disease, perinatal asphyxia, hypoxia, acidosis, hypercapnea.

Source: Maisels MJ, Watchko JF. Treatment of jaundice in low birthweight infants. *Arch Dis Child Fetal Neonatal Ed.* 2003;88:F459-F463.

- Fluorescent tubes
 - Special blue tubes, such as F20 T12/BB, provide more irradiance in the blue spectrum than other tubes and are the most effective fluorescent light source. (2) (Fig. 46.2). The special blue F20 T12/BB tubes provide much greater irradiance than regular blue tubes, labeled F20T12/B. The flickering glare of the blue light has been reported to cause giddiness, nausea, and temporary blurring of vision in nursing personnel (10). One way to overcome this has been to use cool white light in conjunction with the special blue, but this combination can decrease efficacy by as much as 50%, depending on the proportion of cool white light used



FIG. 46.2. Relative spectral content of phototherapy bulbs. Shaded area indicates wavelength effective for phototherapy. Absolute spectral irradiance (μ W/cm²/nm) depends not only on relative power across wavelength of bilirubin absorption but also on total wattage and distance from infant. Although all bulbs provide effective phototherapy for the same wattage, special blue and blue fluorescent bulbs provide the most amount of power in the bilirubin wavelength. (Based on data from Olympic Medical (d); from Warshaw JB, Gagliardi J, Patel A. A comparison of fluorescent and nonfluorescent light sources for phototherapy. Pediatrics. 1980;65:795-798(e); and from Farr PM, Diffey BL. The colour of light for neonatal phototherapy. Arch Dis Child. 1988;63:461-462(f).

- Green/turquoise lamps penetrate the skin to a greater depth, but the advantage over blue light remains debated (12,13 and 14).
- Cool white lamps may be inadequate in sufficiently decreasing total bilirubin levels unless the lights are positioned in close proximity to the infant (11). As mentioned above, this type of light has also been used together with special blue tubes.
- Daylight lamps, like cool white lamps, have a wider wavelength spectrum and are less effective than blue light (11).
- Halogen lamps
 - Halogen spotlight systems utilize single or multiple metal halide lamps as the light source and can provide high irradiance over a small surface area (>20 $\hat{A}\mu W/cm^2/nm$).
 - These units can generate considerable heat, with the potential of causing thermal skin injury; therefore, they must not be in close proximity to the patient.
 - The variable positioning with respect to the distance from the infant as well as heterogeneity of the irradiance can lead to unreliable dosing and unpredictable clinical responses. In addition, they are more expensive than fluorescent bulbs (2).
- Fiber-optic systems
 - UV-filtered light from a tungsten halogen bulb enters a fiber-optic cable and is emitted from the sides and end of fiber-optic fibers inside a plastic pad.
 - The pad emits insignificant levels of heat, so it can be placed in direct contact with the infant to deliver up to $35 \,\mu$ W/cm²/nm of spectral irradiance, mainly in the blue green range (2,15).
 - The orientation of the fiber-optic fibers determines the uniformity of emission and is unique to each of the commercially available devices.
 - The main advantages of these systems are that, while receiving phototherapy, the infant can be held and/or nursed, thereby minimizing infant/parent separation. In addition, covering the infant's eyes is not necessary, preventing further parental anxiety.
 - The main disadvantage of the fiber-optic pads is that they cover a relatively small surface area, and therefore have less efficacy compared to overhead sources. They should not be used as the sole means of providing phototherapy in an infant with significant hyperbilirubinemia (2,3,10).
 - These devices are often used as an adjunct to conventional overhead application of phototherapy to provide "double phototherapy (circumferential phototherapy), which has greater efficacy because greater body surface area is exposed to the light (15).

Two examples of fiber optic systems are

- The Wallaby 3 Phototherapy System (Respironics, Norwell, MA, USA) delivers light at a wavelength of 425 to 475 nm with an average irradiance of 8 to $10 \,\mu$ W/cm²/nm. Two sizes of fiber-optic light panels are available: a 4-in x 5-in neonatal panel and a 3-in x 14-in wrap-around panel, which may be wrapped around the infant.
- The Homed BiliBlanket Phototherapy System (Ohmeda Medical, Laurel, MD, USA) delivers light at a wavelength of 400 to 550 nm; the intensity of light delivered can be controlled, permitting irradiance levels of 15, 25, and 35 μ W/cm²/nm. The fiber-optic panel consists of 2,400 fibers woven into a mat measuring 10 x 20 cm.
- Gallium nitride light-emitting diodes (LEDs)
 - These systems are semiconductor phototherapy devices capable of delivering high spectral irradiance levels of >200 $\hat{A}\mu W/cm^2/nm$ with very little generation of heat within a very narrow emission spectrum in the blue range (460 to 485 nm) (16,17).
 - LEDs have a longer lifetime (>20,000 hours) and have become cost-effective for use in phototherapy devices.
 - An example of an LED system is the neoBLUE System (Natus Medical, San Carlos, CA, USA). This device delivers blue light in the range of 450 to 470 nm with either a low-intensity (12 to 15 $\hat{A}\mu$ W/cm²/nm) or a high-intensity (30 to 35 $\hat{A}\mu$ W/cm²/nm) setting.

D. Technique (Conventional Phototherapy)

Intensive phototherapy is defined as the use of light in the 430- to 490-nm band delivered at 30 μ W/cm²/nm or higher to the greatest body surface area possible (3).

- Position phototherapy unit over infant to obtain desired irradiance (10 to 40 μ W/cm²/nm). Maximal amount of irradiance achieved by standard technique is generally 30 to 50 μ W/cm²/nm. The distance of the light from the infant has a significant effect on the intensity of phototherapy, and to achieve maximal intensity, the lights should be positioned as close as possible to the infant. Fluorescent tubes may be brought within approximately 10 cm of term infants without causing overheating, but halogen spot phototherapy lamps should not be positioned closer to the infant than recommended by the manufacturer, because of the risk of burns (3,18).
- If increased irradiance is required, add additional units or place a fiber-optic phototherapy pad under the infant (15). Additional surface area may be exposed to phototherapy by lining the sides of the bassinet with aluminum foil or a white cloth (19).
- Keep the photoradiometer calibrated and perform periodic checks of phototherapy units to make sure that adequate irradiance is being delivered (3).
- Maintain an intact Plexiglas shield over phototherapy light bulbs in order to block ultraviolet radiation and to protect the infant from accidental bulb breakage.
- Provide ventilation to the phototherapy unit to prevent overheating light bulbs.
- Maintain cleanliness and electrical safety.

E. Technique (Fiber-Optic Phototherapy)

Fiber-optic phototherapy can be used as the sole source of phototherapy or as an adjunct to conventional treatment.

- Insert panel into disposable cover so that it is flat and directed toward infant.
- Place covered panel around infant's back or chest and secure in position. Phototherapy blanket/pad must be positioned directly next to the infant's skin to be effective.

Avoid constriction and skin irritation under the infant's arms if the panel is wrapped around the infant.

- Discard disposable covers after each treatment and when soiled.
- Use eye patches if there is any direct exposure to lights in panel or if used with conventional phototherapy for double-sided effect.
- Ensure stability and adequate ventilation of illuminator unit by placing on a secure surface.
- Connect fiber-optic panel to illuminator.
- Keep fiber-optic panel and illuminator clean and dry.
- Allow lamp to cool for 10 to 20 minutes before moving illuminator. Do not place sharp or heavy objects on panel or cable.

Care of the Infant Receiving Phototherapy

- Monitor temperature, particularly of infants in an incubator, who may develop hyperthermia.
- Monitor intake, output, and weight. Fluid supplementation may be necessary secondary to increased insensible losses and frequent stooling. Encourage breast feeding. Healthy term breast-fed infants may be supplemented with milk-based formula if maternal milk supply is inadequate. Intravenous fluids are rarely required. Milk feeding inhibits the enterohepatic circulation of bilirubin (3).
- The use of eye protection in the form of eye patches is necessary for infants receiving overhead phototherapy. Masks adhering directly to Velcro tabs on the temples are preferable to circumferential headbands.

- Maximize skin exposure to phototherapy source by using the smallest possible diapers as well as keeping blanket rolls from blocking light.
- Avoid fully occlusive dressings, bandages, topical skin ointments, and plastic in direct contact with the infant's skin, to prevent burns.
- Remove plastic heat shields and plastic wrap that decrease irradiance delivered to the skin (20).
- If in use, shield the oxygen saturation monitor probe from the phototherapy light.
- Encourage parents to continue feeding, caring for, and visiting their infant.

F. Home Phototherapy

Home phototherapy decreases costs of hospitalization and eliminates separation of mother and infant. It is safe and effective for selected infants. Home phototherapy should be used only in infants whose bilirubin levels are in the "optional phototherapy range (Fig. 46.1).

- Make arrangements to measure infant's serum bilirubin every 12 to 24 hours, depending on the previous concentration and rate of rise. The infant should be examined daily by a visiting nurse or at an office.
- The supervising physician should be in contact with the family daily during the period of treatment.
- The infant should be rehospitalized if he or she shows signs of illness or if the serum bilirubin concentration exceeds 18 mg/dL.

Efficacy of Phototherapy

The therapeutic efficacy of phototherapy depends on several factors.

- Exposed body surface area: The greater the area exposed, the greater the rate of bilirubin decline
- Distance of the infant from the light source
- Skin thickness and pigmentation
- Total bilirubin at clinical presentation
- Duration of exposure to phototherapy

G. Discontinuation of Phototherapy and Follow-Up

- There is no standard for discontinuing phototherapy. The total serum bilirubin (TSB) level that determines the discontinuation of phototherapy depends on the age at which treatment was initiated and the etiology of hyperbilirubinemia (3,21).
- For infants who are readmitted to hospital (usually for TSB levels of 18 mg/dL or higher), phototherapy may be discontinued when the serum bilirubin level falls below 13 to 14 mg/dL.
- For infants who are readmitted with hyperbilirubinemia and then discharged, significant rebound is uncommon, but may still occur. In cases of prematurity, positive direct antiglobulin (Coombs) test, and for babies treated less than 72 hours, the likelihood of rebound is much higher, and these risk factors should be taken into account when planning postphototherapy follow-up (22,23 and 24). Generally, a follow-up bilirubin measurement within 24 hours after discharge is recommended (3).

H. Complications of Phototherapy

Phototherapy has been used in millions of infants for more than 30 years, and reports of significant toxicity are exceptionally rare(3).

Complications include the following.

- Bronze baby syndrome occurs in some infants with cholestatic jaundice who are exposed to phototherapy, as a result of accumulation in the skin and serum of porphyrins. The bronzing disappears in most infants within 2 months (25). Rare complications of purpuric eruptions due to transient porphyrinemia have been described in infants with severe cholestasis who receive phototherapy (26).
- Diarrhea or loose stools (27)
- Dehydration secondary to insensible water loss
- Skin changes ranging from minor erythema, increased pigmentation, and skin burns to rare and more severe blistering and photosensitivity in infants with porphyria and hemolytic disease
- Potential retinal damage from light exposure if eye patches are not used effectively (28).
- Separation of mother and infant and interference with bonding.

References

1. Vreman HJ, Wong RJ, Stevenson DK. Phototherapy: current methods and future directions. Semin Perinatol. 2004;28:326–333.

2. McDonagh AF, Lighter DA. Phototherapy and the photobiology of bilirubin. Semin Liver Dis. 1988;272–283.

3. American Academy of Pediatrics Subcommittee on Hyperbilirubinemia. Management of

hyperbilirubinemia in the newborn infant 35 or more weeks of gestation. Pediatrics. 2004;114:297–316. 4. Maisels MJ, Watchko JF. Treatment of jaundice in low birthweight infants. Arch Dis Child Fetal Neonatal Ed. 2003:88: F459–F463.

5. Cashore WJ. Bilirubin and jaundice in the micropreemie. Clin Perinatol. 2000;27:171–179.

6. Soylu A, Kavukcu S, Turkmen M. Phototherapy sequela in a child with congenital erythropoietic porphyria. Eur J Pediatr. 1999;158:526–527.

7. Kearns GL, Williams BJ, Timmons OD. Fluorescein phototoxicity in a premature infant. J Pediatr. 1985;107:796–798.

8. Valaez T, Petmezaki S, Henschke C, et al. Control of jaundice in preterm newborns by an inhibitor of bilirubin production: studies with tin-mesoporphyrin. Pediatrics. 1994; 93:1–11.

9. Maisels MJ. Phototherapy: traditional and nontraditional. J Perinatol. 2001(suppl 1):S93–S97.
10. Sarici SU, Alpay F, Unay B, et al. Comparison of the efficacy of conventional special blue light phototherapy and fiberoptic phototherapy in the management of neonatal hyperbilirubinemia. Acta Paediatr. 1999;88:1249–1253.

11. De Carvalho M, De Carvalho D, Trzmielina S, et al. Intensified phototherapy using daylight fluorescent lamps. Acta Pediatr. 1999;88:768–771.

12. Ebbesen F, Agati G, Pratesi R. Phototherapy with turquoise versus blue light. Arch Dis Child Fetal Neonatal Ed. 2003;88: F430–F431.

13. Seidman DS, Moise J, Ergaz Z. A prospective randomised controlled study of phototherapy using blue and blue-green light emitting devices and conventional halogen quartz phototherapy. J Perinatol. 2003;23:123–127.

14. Roll EB, Christensen T. Formation of photoproducts and cytotoxicity of bilirubin irradiated with turquoise and blue phototherapy light. Acta Pediatr. 2005;94:1448–1454.

15. Tan KL. Comparison of the efficacy of fiberoptic and conventional phototherapy for neonatal hyperbilirubinemia. J Pediatr. 1994;125:607–612.

16. Vreman HJ, Wong RJ, Stevenson DK, et al. Light emitting diodes: a novel light source for phototherapy. Pediatr Res. 1998;44: 804–809.

17. Seidman DS, Moise J, Ergaz Z, et al. A new blue light emitting phototherapy device: a prospective randomized controlled study. J Pediatr. 2000;136:771–774.

18. Maisels MJ. Why use homeopathic doses of phototherapy. Pediatrics. 1996;98:283–287.

19. Eggert P, Stick C, Schroder H. On the distribution of irradiation intensity in phototherapy. Measurements of effective irradiance in an incubator. Eur J Pediatr. 1984;142:58–61.

20. Kardson J, Schothorst A, Ruys JH, et al. Plastic blankets and heat shields decrease transmission of phototherapy light. Acta Paediatr Scand. 1986;75:555.

21. Maisels MJ, Kring E. Bilirubin rebound following intensive phototherapy. Arch Pediatr Adolesc Med. 2002;156:669–672.

22. Yetman RJ, Parks DK, Huseby V, Garcia J. Rebound bilirubin levels in infants receiving phototherapy. J Pediatr. 1998;133: 705–707.

23. Lazar L, Litwin A, Merlob P. Phototherapy for neonatal nonhemolytic hyperbilirubinemia. Analysis of rebound and indications for discontinuing phototherapy. Clin Pediatr (Phila.). 1993;32:264–267.

24. Kaplan M, Kaplan E, Hammerman C, et al. Post phototherapy neonatal rebound: a potential cause of significant hyperbilirubinemia. Arch Dis Child. 2006;91:31-–34.

25. Rubaltelli F, Da Riol R, D'Amore ES, et al. The bronze baby syndrome: evidence of increased tissue concentration of copper porphyrins. Acta Pediatr. 1996;85:381–384.

26. Paller AS, Eramo LR, Farrell EE, et al. Purpuric phototherapy induced eruption in transfused neonates: relation to transient porphyrinemia. Pediatrics. 1997;100:360–364.

27. DeCurtis M, Guandalini S, Fasano A, et al. Diarrhea in jaundiced neonates treated with phototherapy: role of intestinal secretion. Arch Dis Child. 1989:64:1161–1164.

28. Bhupathy K, Sethupathy R, Pildes RS. Electroretinography in neonates treated with phototherapy. Pediatrics. 1978;61: 76-81.

47-Intraosseous Infusions

Mary E. Revenis

A. Indications

• Emergency intravenous access when other venous access is not readily available; to restore intravascular volume so that peripheral venous access becomes possible. See Table 47.1 for categories of fluid that have been infused (1,2,3 and 4).

B. Contraindications (3,5,6)

- Bone without cortical integrity (fracture, previous penetration): Extravasation of infusate
- Sternal site: Potential damage to heart and lungs (7)
- Overlying soft tissue infection
- Osteogenesis imperfecta
- Obliterative diseases of marrow such as osteopetrosis

C. Equipment Sterile

- Surgical gloves
- Cup with antiseptic solution
- Gauze squares
- Aperture drape
- 1% lidocaine in 1-mL syringe with 25-gauge needle
- Needle, in order of preference (1,2,3,8)

- Bone marrow or intraosseous needle (18-gauge) (stylet and adjustable depth indicator preferred)
- Short spinal needle with stylet (18- or 20-gauge)
- \circ Short hypodermic needle (18- or 20-gauge)
- Butterfly needle (16- to 19-gauge)
- 5-mL syringe on a three-way stopcock and IV extension set with clamp
- Syringes with saline flush solution
- Intravenous infusion set and intravenous fluid

Nonsterile

- Small sand bag or rolled towel to aid in stabilizing limb
- Tape
- Armboard
- Disposable plastic cup

Optional

Intraosseous needle placement device (intended for use at the proximal tibial location). Devices approved for newborns are the spring-activated B.I.G. Bone Injection Gun (Waismed, Houston, TX, USA) and the battery operated driver EZ-IO Pediatric (Vidacare, San Antonio, TX, USA). Information on use of these devices in premature infants is scarce and as yet unpublished. It is not known if the incidence of success or complications using these devices differs compared to manual insertion of the intraosseous needle.

D. Precautions

- Limit use to emergency vascular access, when peripheral or central venous access is not feasible.
- Avoid inserting needle through infected skin or subcutaneous tissue.
- Stabilize limb with counterpressure, with sand bag or towel roll directly opposite proposed site of penetration, to avoid bone fracture.
- If hand is used to stabilize limb, do not position hand behind puncture site, to avoid inadvertent puncture of hand by the intraosseous needle if it goes through the limb. This is true regardless of whether a sand bag or towel is used. It is at times helpful to use the hand to help stabilize the limb, but it must be done without injury to the practioner.
- Limit needle size to decrease chance of fracture of bone.
- Administer drugs in the usual doses for intravenous administration; however, when possible, dilute hypertonic or strongly alkaline solutions prior to infusion, to reduce risk of bone marrow damage (2).
- Discontinue intraosseous infusion as soon as alternative venous access is established, to reduce risk of osteomyelitis.

E. Technique

Proximal tibia (1,2,3,9,10) (Fig. 47.1)

- Position patient supine.
- Place sand bag or towel roll behind knee to provide countersupport behind puncture site.
- Clean proximal tibia with antiseptic solution.
- Put on sterile gloves.

TABLE 47.1 Types of Intraosseous Infusates Reported in the Literature (4,15,17,21)

- 1. Fluids
 - a. Normal saline
 - b. Crystalloids
 - c. Glucose (dilute if possible when using D50) (15,22)
 - d. Ringer's lactate (17)
- 2. Blood and blood products
- 3. Medications
 - a. Anesthetic agents
 - b. Antibiotics
 - c. Atropine (17)
 - d. Calcium gluconate
 - e. Dexamethasone (17)
 - f. Diazepam (17)
 - g. Diazoxide (17); phenytoin (23)
 - h. Dobutamine (21)
 - i. Dopamine (21,22,24)
 - j. Ephedrine (25)
 - k. Epinephrine (25)
 - l. Heparin (17)
 - m. Insulin
 - n. Isoproterenol (24)
 - o. Lidocaine
 - p. Morphine
 - q. Sodium bicarbonate (dilute if possible) (15,26)
 - Contrast material (27)
- Apply aperture drape.

4.

- If appropriate, inject lidocaine into skin, soft tissue, and periosteum (11).
- Determine penetration depth on needle: Rarely more than 1 cm in infants.
 - For needle or bone needle injection device with adjustable depth indicator, adjust sheath to allow desired penetration.
 - For needle without an adjustable depth indicator, hold the needle in the dominant hand with blunt end supported by the palm and the index finger approximately 1 cm from the bevel of the needle, to avoid pushing it past this mark.
- Palpate tibial tuberosity with index finger.



FIG. 47.1. A: Anterior view. B: Sagittal section. C: Cross section through tibia.

- Hold the thigh and knee above and lateral to the insertion site with the palm of the nondominant hand. Wrap fingers and thumb around the knee to stabilize the proximal tibia.
- Insert needle on the flat, anteromedial surface of the tibia, 1 to 2 cm below the tibial tuberosity.
- Direct needle at an angle of 10 to 15 degrees toward the foot, to avoid the growth plate.

• Advance needle.

- For manual insertion, advance needle using firm pressure with a twisting motion until there is a sudden, slight decrease in resistance, indicating puncture of the cortex.
- If an automatic spring-activated intraosseous needle injection device is used, hold the cylinder against the puncture site at a 90-degree angle with one hand. Release the safety latch on the cylinder with the other hand. Depress the cylinder, as with a syringe, without the use of force.
- If a battery-operated driver with attached needle is utilized, hold the driver in the dominant hand. Position the needle against the puncture site at a 90-degree angle. Depress the trigger to activate the driver. Do not force the driver, but apply firm, steady pressure, allowing the driver to insert the needle. Stop when there is a sudden decrease in resistance.
- Do not advance the needle beyond cortical puncture.
- Remove the stylet.
- Confirm the position of the needle in the marrow cavity.
 - Needle should stand without support.
 - Securely attach a 5-mL syringe and attempt to aspirate blood or marrow. Aspiration is not always successful when using an 18- or 20-gauge needle.
 - If bone marrow is aspirated, it can be analyzed for blood chemistry values, partial pressure of arterial carbon dioxide, pH, hemoglobin level (12,13), type and cross-match, or cultured (13).
 - Attach syringe of saline flush solution and infuse 2 to 3 mL slowly, while palpating the tissue adjacent to the insertion site to detect extravasation. There should be only mild resistance to fluid infusion.
- If marrow cannot be aspirated and significant resistance to fluid infusion is met:
 - The hollow bore needle may be obstructed by small bone plugs.
 - Reintroduce the stylet, or
 - Introduce a smaller-gauge needle through the original needle.
 - Attach syringe of saline flush and flush 2 to 3 mL of fluid.
 - The bevel of the needle may not have penetrated the cortex.
 - Redetermine estimated depth needed.
 - Advance.
 - Flush with saline.
 - The bevel of the needle may be lodged against the opposite cortex.
 - Withdraw needle slightly.
 - Flush with saline.
- Observe the site for extravasation of fluid, indicating that:
 - The placement is too superficial, or
 - The bone has been penetrated completely.
 - If extravasation occurs, withdraw needle and select a different bone.
- When needle position is confirmed:
 - Attach syringe and infuse medications or fluid directly into the needle or via an IV extension set with clamp. Clear medications with saline flush.
 - For continuous infusion, attach a standard intravenous infusion set with an infusion pump to the intraosseous needle and administer at the same rate as for IV infusion (2).
- Secure intraosseous needle and maintain a clean infusion site while the needle is in place.
 - Tape the flanges of the needle to the skin to prevent dislodgement. If a needle safety latch is provided, attach the latch and then apply tape.
 - If desired, cover the exposed end of the needle with a disposable cup, taping the cover down. Cutting off the bottom of the cup will aid in visualization of the site for monitoring.
- Secure intravenous tubing with tape to the leg.
- Secure the leg to the armboard.
- Obtain radiograph to confirm position of needle and to rule out fracture.



FIG. 47.2. Intraosseous infusion into the distal tibia. [Reproduced with permission from Spivey WH. Intraosseous infusions. J Pediatr. 1987;111(5):639.]

• Discontinue intraosseous infusion as soon as alternative intravenous access is achieved.

In an infant with hypotension/hypovolemia, infusion via the interosseous route can restore peripheral perfusion to a point at which venous access is possible in well under 30 minutes.

- \circ Remove needle.
- Apply a sterile dressing over the puncture site.
- Apply pressure to the dressing for 5 minutes.

Distal tibia (2,8) (Fig. 47.2)

- Position patient supine.
- Prepare site and needle as for proximal tibia.
- Insert needle in the medial surface of the distal tibia just proximal to the medial malleolus.
- Direct needle cephalad away from the joint space.
- Proceed as for proximal tibia.

Distal femur (1,3,9) (Fig. 47.1)

- Position patient supine.
- Place sand bag or towel roll behind knee.
- Prepare site and needle as for proximal tibia.
- Insert needle 1 to 3 cm above the external condyles in the anterior midline.
- Direct needle cephalad at an angle of 10 to 15 degrees.
- Proceed as for proximal tibia.

F. Complications (1,4,14,15)

- Fracture of bone (16)
- Complete penetration of bone (17)
- Osteomyelitis (14,15)
- Periostitis (15)
- Subcutaneous abscess
- Cellulitis
- Sepsis
- Extravasation of fluid from the puncture site

- Subperiosteal or subcutaneous infiltration or hematoma
- Compartment syndrome (18)
- Subcutaneous sloughing
- Death (reported only with sternal bone site) (7)
- Theoretical (as yet unreported) (19,20)
 - Embolization of bone fragments or fat
 - $\circ \quad \text{Damage to bone marrow} \\$
 - Damage to growth plate

References

1. Fiser D. Intraosseous infusion. N Engl J Med. 1990;322:1579.

- 2. Spivey W. Intraosseous infusions. J Pediatr. 1987;111:639.
- 3. Hodge D. Intraosseous infusions: review. Pediatr Emerg Care. 1985;1:215.

4. Ellemunter H, Simma B, Trawoger R, Maurer H. Intraosseous lines in preterm and full term neonates. Arch Dis Child Fetal Neonatal Ed. 1999;80:F74.

5. Manley L, Haley K, Dick M. Intraosseous infusion: rapid vascular access for critically ill or injured infants and children. J Emer Nurs. 1988;14:63.

6. Miner WF, Corneli HM, Bolte RG, et al. Prehospital use of intraosseous infusion by paramedics. Pediatr Emerg Care. 1989;5:5.

7. Turkel H. Deaths following sternal puncture. JAMA. 1954;156:992.

8. Iserson K, Criss E. Intraosseous infusions: a usable technique. Am J Emerg Med. 1986;4:540.

9. Carlson DW, DiGuilio GA, Givens TG, et al. Illustrated techniques of pediatric emergency procedures.

In: Fleisher G, Ludwig S, Henretig FM, eds. Textbook of Pediatric Emergency Medicine. 5th ed. Philadelphia: Lippincott Williams & Wilkins; 2006:1879.

10. Parrish G, Turkewitz D, Skiendzielewski J. Intraosseous infusions in the emergency department. Am J Emerg Med. 1986;4:59.

11. Mofenson HC, Tascone A, Caraccio TR. Guidelines for intraosseous infusions. J Emerg Med. 1988;6:143.

 Johnson L, Kissoon N, Fiallos M, et al. Use of intraosseous blood to assess blood chemistries and hemoglobin during cardiopulmonary resuscitation with drug infusions. Crit Care Med. 1999;27:1147.
 Orlowski JP, Porembka DT, Gallagher JM, VanLente F. The bone marrow as a source of laboratory studies. Ann Emerg Med. 1989;18:1348.

14. Rosetti V, Thompson B, Miller J, et al. Intraosseous infusion: an alternative route of pediatric intravascular access. Ann Emerg Med. 1985;14:885.

15. Heinild S, Sondergaard J, Tudvad F. Bone marrow infusions in childhood: experiences from a thousand infusions. J Pediatr. 1947;30:400.

16. La Fleche F, Slepin M, Vargas J, Milzman D. Iatrogenic bilateral tibial fractures after intraosseous infusion attempts in a 3-month-old infant. Ann Emerg Med. 1989;18:1099.

17. Valdes MM. Intraosseous administration in emergencies. Lancet. 1977;1:1235.

18. Vidal R, Kissoon N, Gayle M. Compartment syndrome following intraosseous infusion. Pediatrics. 1993;91:1201.

19. Pediatric Forum. Emergency bone marrow infusions. Am J Dis Child. 1985;139:438.

20. Fiser RT, Walker WM, Seibert JJ, et al. Tibial length following intraosseous infusion: a prospective, radiographic analysis. Pediatr Emerg Care. 1997;13:186.

Berg RA. Emergency infusion of catecholamines into bone marrow. Am J Dis Child. 1984;138:810.
 Neish SR, Macon MG, Moore JW, Graeber GM. Intraosseous infusion of hypertonic glucose and dopamine. Am J Dis Child. 1988;142:878.

23. Walsh-Kelly C, Berens R, Glaeser P, Losek J. Intraosseous infusion of phenytoin. Am J Emerg Med. 1986;4:523.

24. Bilello JF, O'Hair KC, Kirby WC, Moore JW. Intraosseous infusion of dobutamine and isoproterenol. Am J Dis Child. 1991;145:165.

25. Shoor PM, Berrynill RE, Benumof JL. Intraosseous infusion: pressure-flow relationship and pharmacokinetics. J Trauma. 1979;19:772.

26. Spivey WH. Comparison of intraosseous, central and peripheral routes of sodium bicarbonate administration during CPR in pigs. Ann Emerg Med. 1985;14:1135.

27. Cambray EJ, Donaldson JS, Shore RM. Intraosseous contrast infusion: efficacy and associated findings. Pediatr Radiol. 1997;27:892.

48-Tapping Ventricular Reservoirs

Secelela Malecela

The subcutaneous ventricular access device (Fig. 48.1) is used as a temporary measure to drain cerebrospinal fluid (CSF) in preterm infants with posthemorrhagic hydrocephalus while awaiting permanent ventriculoperitoneal (VP) shunt placement (1,2,3,4,5,6,7).

A ventricular reservoir is often inserted in preterm infants who are too small or too unstable to have a VP shunt. A ventricular reservoir also allows for drainage and clearing of the initial CSF, which is bloody and has a high protein content, thus decreasing the risk of shunt blockage (5,6 and 7).

The reservoir is usually tapped immediately following placement by the neurosurgeon to ensure proper placement and to drain excess CSF (5). Subsequent taps are performed in the neonatal intensive care unit (NICU), aiming to remove enough CSF to prevent further ventriculomegaly and maintain normal head growth (5,6,8).

A. Indications

- Rapidly increasing head circumference, more than 1 cm/week
- Clinical signs of raised intracranial pressure evidenced by a full or tense anterior fontanelle, separation of the sutures, apnea and bradycardia, poor feeding and vomiting
- Ultrasound or radiologic evidence of progressive ventriculomegaly

B. Contraindications

- Low circulating blood volume
- Cellulitis or abrasion over the reservoir site
- Sunken fontanelle or overlapping sutures
- Severe coagulopathy

C. Equipment

- Mask, cap, sterile gloves
- Standard infant lumbar puncture set
- Povidine–iodine surgical scrub and prep solution
- Aperture drape
- Scalp-vein needle (25- or 27-gauge)
- 20-mL syringe

D. Precautions

- Use strict aseptic technique.
- Maintain continuous cardiorespiratory monitoring during the procedure.
- Do not use local anesthetic.
- Do not place intravenous lines on the same side of the scalp.
- Do not shave the operative area.
- Always use a fresh site for insertion of the needle with every tap.
- Avoid puncturing the bottom of the reservoir.

E. Technique

- Infant should be restrained and comfortable, with the head in neutral position.
- Clip any long hair that interferes with the surgical area but do not shave the operative area.
- Clean skin over the reservoir and a radius of 5 cm of the surrounding skin using surgical scrub. Use light but firm contact. Clean for 5 to 10 min.
- Dry with blotting pads.
- Don mask and cap.
- Scrub hands and put on sterile gloves.
- Paint area with povidine–iodine solution and allow area to dry.
- Drape area while maintaining patient visibility.
- Insert scalp-vein needle at an angle of 30 to 45 degrees through the skin into the reservoir bladder (Fig. 48.2).
- Aspirate fluid at a rate of 1 to 2 mL/min. Remove no more than 10 to 15 mL/kg. Some authors advocate letting the CSF drain spontaneously, rather than aspirating, in order to reduce fresh bleeding into the ventricles (9).
- Remove needle and hold firm pressure for 2 minutes, until CSF leakage from the skin stops.
- Clean area with sterile saline to remove the povidine–iodine.
- Remove the restraints.
- Send CSF sample for culture, cell count, glucose, and protein. This should be done every 3 days. If fluid is dark and bloody, it is reasonable to send only a culture sample.



FIG. 48.1. McComb reservoir. Ventricular access device: side (top) and front views (bottom).

- F. A Successful Tap
 - At the end of the procedure the anterior fontanelle should be soft and flat (not sunken) and the cranial bones should be approximated well at the sutures.
 - If sufficient volume is removed, the fontanelle may be full 24 hours later, but the sutures should not split apart.
 - If the fontanelle remains slack, the interval for tapping may be lengthened to every other day and/or the amount of CSF removed at each tap reduced.



FIG. 48.2. Tapping of the reservoir.

	1 Complications of Ventricalar Reservoir Dramage
Problem (Incidence)	What To Do
Hyponatremia (20%–60%)	Monitor serum electrolytes every other day and supplement sodium intake.
Hypoproteinemia (15%)	Ensure adequate protein intake. Monitor serum albumin weekly.
Infection (0%–8%)	A combination of intravenous and intrareservoir antibiotics may rarely be successful. Removal of the reservoir is usually necessary.
Subgaleal CSF collection	Percutaneous aspiration of fluid using a different needle at the same time
(0%a€ 9%)	Tap larger volume of CSF from the reservoir or increase frequency of taps to reduce pressure.
CSF leaks through incision (0%–3%)	Increase frequency of reservoir taps.
Ventricular access device occlusion (0%–10%)	Replace reservoir.
Trapped contralateral ventricle (6%)	Place second reservoir.
Fresh bleeding into the ventricle (0%–40%)	Prevent by using 25- or 27-gauge needle, aspirate slowly or let CSF drain spontaneously rather than aspirating.
Bradycardia, pallor, hypotension (rare)	Stop aspiration. Infuse 10–15 mL/kg of normal saline IV rapidly. Remove a smaller volume at a slower rate at next tap.
Skin breakdown over reservoir (rare)	Avoid abraded skin when tapping the reservoir. Avoid excoriating skin while prepping site.

TABLE 48.1 Complications of Ventricular Reservoir Drainage

G. Follow-Up

- Assess clinical response to taps, daily head circumference, and weekly cranial ultrasonography.
- Interval between taps may range from twice a day to once every 2 to 3 days.
- Taps should be continued until the infant weighs 2 kg and is a suitable candidate for shunt placement or until the hydrocephalus resolves.

H. Complications

See Table 48.1 (1,2,3,4,5,6,7,8,9,10).

References

1. Whitelaw A, Cherian S, Thoresen M, Pople I. Posthaemorrhagic ventricular dilatation: new mechanisms and new treatment. Acta Paediatr. 2004;44:11–14.

2. Benzel EC, Reeves JP, Nguyen PK, Hadden TA. The treatment of hydrocephalus in preterm infants with intraventricular hemorrhage. Acta Neurochir. 1993;122(3–4):200–203.

3. De Vries LS, Liem KD, Van Dijk K, et al.; Dutch Working Group of Neonatal Neurology. Early versus late treatment of posthemmorhagic ventricular dilatation: results of a retrospective study from five neonatal intensive care units in The Netherlands. Acta Paediatr. 2002;91(2):212–217.

4. Heep A, Engelskirchen R, Holschneidr A, Groneck P. Primary intervention for posthemorrhagic hydrocephalus in very low birth weight infants by ventriculostomy. Childs Nerv Syst. $2001;17(1\hat{a}\in 2):47\hat{a}\in 51$.

5. Firm DM, Scott RM, Madsen JR. Surgical management of neonatal hydrocephalus. (Review). Neurosurg Clin North Am. 1998;9(1):105–110.

6. McComb JG. Management of intraventricular hemorrhage in neonates. Western J Med. 1984;769–770.
7. Gaskill SJ, Marlin AE, Rivera S. The subcutaneous ventricular reservoir: an effective treatment for posthemorrhagic hydrocephalus. Childs Nerv Syst. 1988;4(5):291–295.

8. Hudgins RJ, Boydston WR, Gilreath CL. Treatment of posthemorrhagic hydrocephalus in the preterm infant with a ventricular access device. Pediatr Neurosurg. 1998;29(6): 309–313.

9. Moghal NE, Quinn MW, Levene MI, et al. Intraventricular hemorrhage after aspiration of ventricular reservoirs. Arch Dis Child. 1992;67:448–449.

10. Weparin BE, Swift DM. Complications of ventricular shunts. Techn Neurosurg. 2002;7(3):224–242.

49-Laser for Retinopathy of Prematurity

William F. Deegan Marijean Miller

A. Classification of Retinopathy of Prematurity (ROP) (1)

- Location (Fig. 49.1)
 - Zone I: Circle whose center is the optic disc and whose radius is twice the distance from the optic disc to the center of the macula
 - Zone II: Circle whose radius extends from the optic disc to the nasal ora serrata and is peripheral to Zone I
 - Zone III: Temporal crescent of retina anterior to Zone II

• Extent of disease

The retina is divided into 12 equal segments, or clock hours. The extent of retinopathy specifies the number of clock hours involved.

- Staging the disease (1,2) (Figs. 49.2 and 49.3)
 - Stage 1 demarcation line: A flat white line in the plane of the retina, separating avascular retina anteriorly from vascularized retina posteriorly
 - Stage 2 ridge: Elevated fibrovascular tissue extending out of the plane of the retina and separating the vascularized and avascular retina
 - Stage 3 neovascularization of the ridge: Neovascular tissue extending from the ridge into the vitreous. This tissue may cause the ridge to appear ragged or "fuzzy†(Fig. 49.2).
 - Stage 4 retinal detachment: A separation of the retina from the underlying choroid. Usually, traction by the vitreous, through the presence of neovascular tissue, pulls the retina away from its underlying attachments. The intervening (subretinal) space fills with a proteinaceous fluid. A stage 4A detachment spares the macula; a stage 4B involves the macula. Retinal detachments in ROP require more extensive intervention, i.e., incisional surgery (scleral buckling and/or vitrectomy) (3).
 - Stage 5 total retinal detachment: Retinal tissue becomes inextricably bound to reactive vitreous and is pulled by the vitreous into the retrolental space (hence the older term, retrolental fibroplasia).
 - Plus disease: Dilation and tortuosity of retinal vessels due to high-flow shunting in advanced ROP. This is seen best in the posterior pole.
 - Threshold ROP: Traditionally (CRYO-ROP) defined as 5 contiguous or 8 total clock hours of stage 3 with Plus in Zone(s) I and/or II. Any stage 3 or any Plus in Zone I ROP should be treated. The Early Treatment for Retinopathy of Prematurity (ET-ROP)

Study (4) devised a risk score that further stratifies $\hat{a} \in \hat{c}$ threshold $\hat{a} \in \hat{c}$ and suggests earlier, $\hat{a} \in \hat{c}$ treatment in some infants. These include babies with stage $2\hat{a} \in 3$ with Plus disease in Zone II.



FIG. 49.1. Scheme of retina of right eye (RE) and left eye (LE), showing zone borders and clock hours employed to describe location and extent of retinopathy of prematurity. (From Committee for Classification of Retinopathy of Prematurity. An international classification of retinopathy of prematurity. Arch Ophthalmol. 1984;102:1130, with permission.)



FIG. 49.2. Dilated, tortuous vessels end in vascular shunts at a thickened ridge of fibrovascular tissue. Avascular retina lies anterior to the ridge.

• Additional features: Iris vascular engorgement (Fig. 49.3) and pupillary rigidity (manifested by poor dilation after mydriatic instillation) (5) are harbingers of active, advanced ROP; corneal and lenticular opacity can be seen in the eyes of any premature infant regardless of the presence of ROP.



FIG. 49.3. Dilation and tortuosity of iris vessels may be seen in severe threshold retinopathy of prematurity.

B. Screening for Retinopathy of Prematurity (2)

Indirect ophthalmoscopy with scleral depression is the only method that can accurately screen and monitor babies with ROP. All babies born at <1,500 g and/or 32 weeks' or less gestational age should be screened, and infants weighing >1,500 g or more than 32 weeks' gestation should be considered for screening if they have had a particularly unstable clinical course, e.g., those requiring cardiopulmonary support. The exams are done at the bedside with the assistance of the baby's nurse.

The timing of the first exam varies with gestational age. The initial examination for infants born between 22 and 27 weeks' gestational age is at 31 weeks postconceptional age (gestational age at birth plus chronological age). Infants born later than 27 weeks' should be screened initially 4 weeks after birth. Follow-up exams depend on the retinal findings as classified by the International Classification of ROP. Table 49.1 has been adapted from the joint policy statement of the American Academies of Pediatrics and Ophthalmology, and the American Association for Pediatric Ophthalmology and Strabismus (2). Babies whose clinical condition deteriorates should be followed closely, i.e., weekly, as late reactivation and worsening are possible.

C. Indications for Laser (2,4,6)

- The guidelines for consideration of ablative treatment for ROP have been revised recently, according to the Early Treatment for Retinopathy of Prematurity Study.
 - Zone I: Any stage of ROP with Plus disease or stage 3 without Plus disease
 - Zone II: Stage 2 or 3 ROP with Plus disease
- Babies who meet the traditional definition of threshold in Zones II and III according to the Multicenter Trial of Cryotherapy for Retinopathy of Prematurity (CRYO-ROP): stage 3 ROP in Zone I or II in 5 or more continuous clock hours or 8 cumulative clock hours, with Plus disease
- Treatment is recommended within 72 hours of detection of a stage of ROP requiring ablative therapy, when possible, in order to minimize the risk of retinal detachment.

TABLE 49.1 Follow-up Examination Schedule (2)

Findings	Follow-up
Stage 1-2 in Zone 1	1 wk or less
Stage 3 in Zone II	
Immature retina (no ROP) in Zone I	1-2 wk
Stage 2 in Zone II	
Regressing ROP in Zone I	
Stage 1 in Zone II	2 wk
Regressing ROP in Zone I	
Immature retina (no ROP) in Zone I	I2-3 wk
Regressing or Stage 1-2 in Zone III	
D. Contraindications	

- Stage 4 to 5 ROP, in which case laser may be done (intraoperatively) in conjunction with incisional surgery (scleral buckle, vitrectomy, or both)
- Vitreous hemorrhage sufficient to obscure a view of the retina
- Instability of medical condition sufficient to make the stress of sedation and laser inadvisable
- Lethal medical illness

E. Laser Treatment

Personnel

- Ophthalmologist
 - Determines the need for treatment
 - Administers topical anesthetic
 - Ensures that all personnel present at the treatment are wearing laser safety goggles
 - Performs the laser
 - Watches for and treats ocular complications that may arise during and after the procedure
 - Follows the baby postoperatively until ROP is resolved
- Neonatology fellow, attending neonatologist, or pediatric anesthesiologist

- Administers systemic sedative agents (midazolam, fentanyl, ketamine, or a combination)
- Monitors patient for and treats any systemic complications that develop during or after treatment
- Provides information to the ophthalmologist regarding the patient's overall condition throughout the procedure
- Assistant to the ophthalmologist
 - Helps with laser and instruments
 - Records the treatment parameters used during treatment
- Neonatal nurse
 - Instills dilating drops several times in the hour preceding treatment
 - Immobilizes the patient during treatment
 - Monitors the patient's airway

Equipment

- Cardiorespiratory, blood pressure, and transcutaneous oxygen monitors
- Appropriate respiratory support (ventilator, laryngoscope and endotracheal tubes, face mask, selfinflating resuscitation bag, suction, and oxygen source)
- Emergency medications (atropine, epinephrine, bicarbonate, calcium, phenobarbital)

Note: Precalculation of weight-appropriate doses is helpful.

- Topical ocular anesthetic, e.g., tetracaine, proparacaine
- Cycloplegic/mydriatic eye drops: Cyclomydril (Alcon Laboratories, Fort Worth, TX, USA) (cyclopentolate hydrochloride 0.2% and phenylephrine hydrochloride 1%) or 0.5% cyclopentolate and 1% or 2.5% phenylephrine
- Calcium alginate-tipped nasopharyngeal applicators or Flynn depressor (Fig. 49.4), for scleral depression
- Balanced salt solution for rewetting cornea during procedure
- Neonatal eyelid speculum (Fig. 49.4)
- 28- and 20-diopter lenses
- Portable argon or diode laser (7,8) with indirect (headlamp) delivery system
- Appropriate laser safety goggles



FIG. 49.4. Lid speculae and Flynn depressor.
- F. Precautions and Complications (Table 49.2)
 - Ensure that laser is fully functional.
 - If the infant is at high risk for an adverse event that would terminate treatment prematurely, treat the more advanced eye first (assuming both have threshold ROP).
 - Discontinue feedings at least 4 hours before the procedure, or empty the stomach with an orogastric tube.
 - Establish intravenous access for infusions of medications and intravenous fluids.
 - Observe transcutaneous or saturation monitor carefully, and adjust administered oxygen appropriately.
 - Stabilize the infant: Correct electrolyte imbalances, platelet deficiency, etc.
 - Use only 1% phenylephrine if there is a history of hypertension.

Complication	Treatment/Action		
Systemic: intra- and immediately postop			
Bradycardia	Interrupt treatment.		
	Assess airway, oxygen delivery.		
	Atropine 0.1 mg IV		
Hypoxia/cyanosis	Evaluate airway.		
	Administer supplemental oxygen.		
Apnea	Evaluate airway.		
	Gentle stimulation.		
	Administer supplemental oxygen.		
	Hand-ventilate (self-inflating resuscitation bag, face		
	mask).		
Tachycardia	Assess pain control.		
<i>y</i>	Administer additional analgesic.		
	Monitor blood pressure and perfusion.		
Hypertension	Assess pain control.		
	Administer additional analgesic.		
	If moderate, observe.		
	If severe, consider hydralazine 0.1 mg/kg IV		
Arrhythmia	Manage as appropriate for arrhythmia.		
Seizure (mechanism uncertain:?	Supportive care		
anticholinergic effect)	Phenobarbital		
Ocular: intraop	i nonoourorum		
Closure of central artery	Relieve pressure on globe (stop scleral depression).		
Corneal clouding/abrasion	Rinse with balanced salt solution/saline		
Comour crouding, doruston	Interrupt treatment		
Retinal/vitreous/choroidal hemorrhage	Gentle pressure on globe (until arterial pulsations		
Retinal (Recoust encrored nemorinage	visible)		
	Avoid lasering blood		
	May have to terminate treatment if extensive		
Ocular: postop	way have to terminate treatment if extensive.		
Conjuntival hemorrhage	Observation		
Conjunctival laceration	Antibiotic ointment t i d for 33€"4 d		
Corneal abrasion	Antibiotic ointment t i d for $33 \in 4$ d		
Corrical abrasion	Follow with slit lamp evan with fluorescein		
Hynhema	Topical cycloplegic and steroids		
Tryphema	Follow intraocular pressure closely		
	Consider washout if high pressure no resolution in 7-		
	10 d		
Retinal/vitreous/choroidal hemorrhage	Close follow-up		
Ocular: lata	Close lollow-up.		
Amblyonia strabismus myonia	Pediatric onthe Inclose assessment 22Et4 me after		
Amoryopia, suaoisinus, myopia	treatment(s)		
	uvaulutill(5). Educate parents prior to discharge revised for regular		
	and the parents prior to discharge re: need for regular		
	ophulalilology lollow-up.		

TABLE 49.2 Complications of Laser for Retinopathy Treatment/Action

IV, intravenously; t.i.d., three times per day.



FIG. 49.5. Freshly lasered avascular retina.

- Wipe off any excess drops spilling onto the skin to avoid transcutaneous absorption (skin vessel blanching occurs with phenylephrine).
- G. Technique
 - General preparation
 - Instill eyedrops (per orders from ophthalmologist) into both eyes in the hour prior to procedure. Maximal dilation is critical for optimum laser; therefore, several (three or four) instillations of drops may be required, especially in eyes with neovascularization/vascular engorgement of the iris.
 - Transport the patient to surgical suite or designated procedure room in the nursery.
 - Ensure monitors are attached and functioning.
 - Immobilize infant: Swaddle in a clean towel or blanket to immobilize arms and legs.
 - Ensure that the intravenous tubing is accessible.
 - Administer intravenous sedation.

If local anesthesia is to be used, a combination of topical (e.g., tetracaine, proparacaine) and systemic analgesic/sedative e.g. intravenous morphine, medications is administered prior to injection.

- Distribute laser safety goggles and dim overhead lights.
- Retract lids.
- Perform laser: Cover the avascular retina with confluent gray-white burns (Fig. 49.5).
- Have an assistant count and record the number of spots and the duration and power of each spot.

H. Postoperative Care

- Instill 0.25% scopolamine hydrobromide in treated eye(s) daily for 3 to 5 days.
- Apply antibiotic–steroid preparation (e.g., tobramycin–dexamethasone) to treated eye(s) three to four times daily for 5 to 7 days.
- Monitor the patient with a cardiorespiratory monitor for 24 to 72 hours.
- Perform a dilated retinal exam 1 to 2 weeks after treatment.
- If opaque media are present at the time of laser, or if the pupil does not dilate adequately, complete treatment of the avascular retina may be impossible, and "skip areas†may be visible in the

weeks after treatment. Treatment of these areas should be considered if there is not marked resolution of the adjacent plus disease and/or neovascularization.

• Follow the infant every 1 to 2 weeks until the ROP resolves completely. If at the time of discharge ROP is still present, ensure that the parents and the physicians responsible for the care of the infant after discharge are aware of the extreme importance of maintaining a regular schedule of outpatient examinations. Once the ROP has resolved completely, the baby should be seen by a pediatric ophthalmologist within 1 to 2 months to assess vision, ocular alignment and motility, refractive status, etc. Long-term follow-up over several years may be necessary (9,10,11 and 12).

I. Postdischarge Care

A critical component of treatment is post-discharge care. It is imperative that infants who develop any stage of ROP, especially those with prethreshold stage 3 or those that have received treatment, are seen within 1 to 2 weeks of discharge, or as directed by the ophthalmologist involved in the baby's care. No baby with any ROP, or who has regressed ROP after treatment, should leave the neonatal intensive care unit (NICU) without a scheduled follow-up examination.

A careful, reproducible tracking system for arranging follow-up should be established by every NICU. A member of the staff of each NICU should be responsible for maintaining and periodically auditing this system.

Verbal and written instructions for follow-up should be given to the parents. Parents should be given a discharge form indicating their baby's scheduled follow-up among their discharge instructions. The importance of scheduled follow-up should be prominently stated on the form.

J. Outcome

The benefit of ablative therapy for threshold ROP became apparent so quickly that enrollment for the CRYO-ROP study was halted after approximately half the planned number of infants were enrolled. The risk of adverse outcome (retinal detachment, macular fold) was cut in half in treated eyes. Subsequently, as indirect argon and diode laser became available in the mid-1990s, the risk of adverse outcome in Zone II and III disease was reduced by another 50% (7,10). The outcome for eyes with Zone I disease, although poor, has improved with laser and incisional surgery (vitrectomy). Specifically, laser treatment of the posterior avascular retina can be accomplished easily and without necessitating conjuctival incisions, as in cryotherapy.

Treated eyes carry a risk of retinal dystopia, and myopia, and subsequent strabismus and amblyopia (11,12 and 13). To minimize the effect of refractive errors and strabismus, careful follow-up by a pediatric ophthalmologist is mandatory.

Premature infants are at risk for intracranial pathologies that may limit visual function. Pediatric ophthalmologists, neurologists, and others involved in the care of former preemies should be in frequent contact in order to address the often complex and changing visual deficits present in these children.

References

1. Committee for Classification of Retinopathy of Prematurity. An international classification of retinopathy of prematurity. Arch Ophthalmol. 1984;102:1130.

2. Section on Ophthalmology, American Academy of Pediatrics, American Academy of Ophthalmology, American Association for Pediatric Ophthalmology and Strabismus. Screening examinations of premature infants for retinopathy of prematurity. Pediatrics. 2006;117:572–576.

3. Trese MT. Scleral buckling for retinopathy of prematurity. Ophthalmology. 1994;101:23–26.
4. Early Treatment for ROP Cooperative Group. Revised indications for the treatment of retinopathy of prematurity: results of the early treatment for retinopathy of prematurity randomized trial. Arch Ophthalmol. 2003;121:1684–1694.

5. Kivlin JD, Biglan AW, Gordon RA, et al., for Cryotherapy for Retinopathy of Prematurity Group. Early retinal vessel development and iris vessel dilatation as factors in retinopathy of prematurity. Arch Ophthalmol. 1996;114:150–154.

6. Multicenter Trial of Cryotherapy for Retinopathy of Prematurity Cooperative Group. Manual of Procedures. PB 88-163530. Springfield, VA: National Technical Information Service, U.S. Department of Commerce; 1985.

7. Laser ROP Study Group. Laser therapy for retinopathy of prematurity. Arch Ophthalmol. 1994;112:154–156.

8. Hunter DG, Repka MX. Diode laser photocoagulation for threshold retinopathy of prematurity. Ophthalmology. 1993;100:238–244.

9. Cryotherapy for Retinopathy of Prematurity Cooperative Group. Multicenter trial of cryotherapy for retinopathy of prematurity: one-year outcomeâ€"structure and function. Arch Ophthalmol. 1990;108:1408â€"1416.

10. McNamara J. Laser therapy for retinopathy of prematurity. Curr Opin Ophthalmol. 1993;4:76–80. 11. O'Connor AR, Stephenson T, Johnson A, et al. Long-term ophthalmic outcome of low birth weight children with and without retinopathy of prematurity. Pediatrics. 2002;109:12–18.

12. Larson EK, Rydberg AC, Holmstrom GE. A population-based study of the refractive outcome in 10year-old pre-term and full-term children. Arch Ophthalmol. 2003;121:1430–1436.

13. Gilbert WS, Dobson V, Quinn GE, et al., on behalf of the Cryotherapy for Retinopathy of Prematurity Cooperative Group. The correlation of visual function with posterior retinal structure in severe retinopathy of prematurity. Arch Ophthalmol. 1992;110:625–631.

50-Peritoneal Dialysis

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Acute Peritoneal Dialysis (1,2,3,4, and 5)

In neonates, acute peritoneal dialysis (PD) is frequently preferred over hemodialysis (HD), continuous arteriovenous hemofiltration with or without dialysis (CAVH/D), and continuous venovenous hemofiltration with or without dialysis (CVVH/D) because it is technically easier. Because peritoneal surface area per kilogram of body weight is relatively larger in newborns and children than in adults, peritoneal dialysis usually allows adequate clearance and removal of excess fluid (6); in addition, it avoids the need for anticoagulation and maintenance of adequate vascular access (7).

A. Indications

- Renal failure, when conservative management has failed to adequately control any of the following conditions (8,9):
 - o Hypervolemia
 - o Hyperkalemia
 - Hyponatremia
 - Refractory metabolic acidosis
 - Hyperphosphatemia
 - Azotemia
 - Additional fluid space needed for delivering drugs and/or nutrition
- Inherited disorders of organic and amino acid metabolism when HD or CVVH/D is unavailable (10,11)

- In hyperammonemic metabolic crisis, evidence suggests that ammonia is more efficiently removed by extracorporeal techniques than by PD (12).
- In babies with imminent or current intracranial hemorrhage, PD is considered the therapeutic option of choice, especially in nonhyperammonemic disorders (12).
- **B.** Relative Contraindications
 - Acute abdomen
 - Abdominal adhesions
 - Immediately after abdominal surgery (13)
 - Diaphragmatic or abdominal wall disruptions
- C. Equipment (Figs. 50.1, 50.2 and 50.3)
 - Masks, sterile drapes, gowns, and gloves
 - Povidone–iodine
 - 1% xylocaine without epinephrine
 - 3-mL syringe with 25-gauge needle
 - Intravenous cutdown tray with no. 11 surgical blade
 - Waterproof tape
 - 22-gauge angiocatheter
 - A temporary catheter such as a 14-gauge angiocatheter or one of the commercially available temporary dialysis catheters, e.g., a Trocath (Trocath Peritoneal Dialysis Center, Kendall McGaw Laboratories, Sabana Grande, Puerto Rico)
 - Dialysis solution (1.5, 2.5, or 4.25%)
 - Other concentrations can be made by manual mixing of standard solutions
 - Heparin
 - Home Choice Automated PD System (Fig. 50.2) or any other reliable fluid warmer
 - IV pole
 - Inline burette set
 - Ultra Set CAPD Disposable Disconnect Y-Set
 - MiniCap Extended Life PD Transfer Set With Twist Cap
 - FlexiCap Disconnect Cap with povidone–iodine solution
 - Medicap with povidone–iodine solution
 - Baby weigh scale with low resolution (Medela, which has a resolution of 2 g from 0 to 6,000 g) (Fig. 50.2)

An alternative approach is to utilize a pediatric cycler set. Experience in using this equipment is necessary. We recommend a commercially available cycler that provides a minimum fill volume of 50 mL with 10-mL increments.



FIG. 50.1. A, IV pole (Fig. 50.3); B, Dianeal PD-2 Peritoneal Dialysis Solution (Baxter Healthcare Corporation, Deerfield, IL, USA); C, Inline Burette Set 150 mL (Abbott Laboratories, North Chicago, IL, USA); D, Ultraset CAPD Disposable Disconnect Y-Set (Baxter Healthcare Corporation, Deerfield, IL, USA); E, MiniCap Extended Life PD Transfer Set With Twist Clamp (Baxter Healthcare Corporation, Deerfield, IL, USA); F, Flexicap Disconnect Cap with Povidone–Iodine Solution (Baxter Healthcare Corporation, Deerfield, IL, USA); G, Minicap with Povidone–Iodine Solution (Baxter Healthcare Corporation, Deerfield, IL, USA).

D. Preprocedure Care

- Obtain informed consent.
- Check body weight and abdominal girth.
- Check for infection at the insertion site.
- Decompress the stomach.
- Catheterize the bladder.
- Place preweighed diaper under the patient.

Before assembly of system, wash hands and put on a mask. All connections should be made using sterile technique. Universal precautions should be observed (Chapter 4). Keep all tubing clamped. See Fig. 50.3 for connections.



FIG. 50.2. Right: Home Choice Automated PD System (Baxter Healthcare Corporation, Deerfield, IL, USA). Left: Medela baby weigh scale (Medela, McHenry, IL, USA).



FIG. 50.3. An assembled peritoneal dialysis circuit illustrates an IV pole (A) and an in-line burette (C) that is connected to an Ultra Set CAPD Y-Set (D). The short limb of this Y-Set is connected to the transfer set (E), which is connected to a Tenckhoff catheter exiting from the abdominal cavity of a doll, and its long limb has a bag at its end that is located on the floor.

- Add 500 U of heparin to each 1 L of the dialysis solution. Start with 1.5% dialysate.
- Warm a liter bag of dialysate (Dianeal or other), or a larger bag if 1 L dialysate is not unavailable, by resting it on the heating surface of the Home Choice Automated PD System, or a reliable fluid

warmer. The temperature can be set between $35\hat{A}^{\circ}C$ and $37\hat{A}^{\circ}C$. For a newborn, keep the temperature at $37\hat{A}^{\circ}C$ (in older pediatric patients, the temperature is usually set to $36\hat{A}^{\circ}C$, and occasionally to $35\hat{A}^{\circ}C$ if the environmental temperature is high).

- Spike the in-line burette set (Abbott Laboratories, North Chicago, IL, USA) into the dialysate (Dianeal or other) when the ideal temperature has been achieved.
- Connect the outlet of the burette set to the inflow line of the Ultra Set CAPD Disposable Disconnect Y-Set.
- Connect the short arm end of the Y-Set to the twist clamp end of a MiniCap Extended Life PD Transfer Set with Twist Clamp (Baxter Healthcare Corporation, Deerfield, IL, USA). If the catheter is placed surgically, this Transfer Set is routinely connected by most surgeons to the Tenckhoff catheter, before assessment of patency, and you will be able to skip this step.
- Prime the circuit in a sterile fashion, clamp, and cap the end of the transfer set, or the short limb of the Y-Set.

E. Procedure (Also Refer to Chapter 25, Abdominal Paracentesis, and Videos on Procedures DVD.)

The ideal technique is surgical insertion of a permanent peritoneal dialysis catheter, which can be placed in the neonatal intensive care unit (14). The catheter is tunneled from the peritoneum to an exit site on the skin; it usually works well and leaks infrequently (Quinton Pediatric Tenckhoff Neonatal 31 cm² Cuff catheter. Kendall Healthcare, Mansfield, MA, USA). However, if surgical insertion of a permanent catheter is not possible, an alternative approach is to utilize an angiocatheter or a temporary PD catheter for no longer than a few days. Note that surgically inserted catheters are associated with fewer acute complications (15). With catheters inserted at the bedside, guidewire-inserted femoral catheters have shown the least mechanical complications; intravenous catheters produce more mechanical complications than femoral catheters, but less than stylet catheters (16,17).

- Monitor vital signs.
- Restrain infant in supine position.
- Scrub.
- Prepare the skin of the abdomen (Chapter 4).
- Drape to expose the insertion site.

The choice of insertion site is influenced by the preference of the physician and/or the presence of postoperative wounds, abdominal wall infection, or organomegaly. A location one third the distance from the umbilicus to the symphysis pubis in the midline or a site lateral to the rectus sheath in either of the lower quadrants is preferred.

- Infuse approximately 0.5 mL of xylocaine around the insertion point.
- Select either a 14-gauge angiocatheter or a temporary dialysis catheter.
- If you elect to use a 14-gauge angiocatheter:
 - Insert the angiocatheter at the insertion site.
 - Remove the stylet.
 - Infuse ~20 mL of normal saline to confirm a free flow. Clamp.
 - Proceed to Step 10 (Section E).
- If using a soft and flexible temporary catheter, such as a Cook catheter (Cook Critical Care, Bloomington, IN, USA), follow the manufacturer's instructions. Then proceed to Step 10 (Section E).
- Test patency.
 - Temporary catheter:
 - May observe flow of a few drops of saline. Connect the free end of the Transfer Set to the catheter.
 - Allow approximately 30 mL of dialysis solution to enter peritoneal cavity by gravity.

- Clamp the short arm of the Y-Set (inflow).
- Unclamp the long arm of the Y-Set (outflow).
- Repeat steps a(2) through a(4) several times.
- Secure the temporary catheter with a purse-string suture and tape if inflow and outflow occur readily.
- Tenkhoff catheter:
 - Unclamp the Transfer Set. Will observe either saline or dialysis fluid, which was instilled at surgery, draining. Allow to drain to completion. Connect the short arm of the Y-Set to the Transfer Set.
 - Follow steps a(2) through a(5) of Step 10 above.

This procedure (Step 10) usually results in a positive fluid balance (the volume drained is less than the volume infused). This retention is acceptable.

F. Management

- Establish a cycle time. This is usually about 60 minutes and consists of a fill by gravity, dwell time of 45 minutes, and drain by gravity.
- Establish a dialysis volume per pass. Starting volume is usually 20 to 30 mL/kg.
- Clamp the long arm of the Y-Set (outflow line).
- Unclamp the inflow line.
- Allow the dialysate to flow in as quickly as possible while carefully observing vital signs.
- Clamp the inflow line.
- Allow the fluid to dwell.
- Unclamp the outflow when dwell time is completed.
- Allow 5 to 10 minutes for draining.
- Clamp the outflow line.
- Repeat the cycle.
- Increase the volume by 5 mL/kg/cycle slowly. Maximum volume is 40 mL/kg if tolerated, attained over 12 to 24 hours.
- Continue to add 500 U of heparin/L of dialysate, until dialysate effluent return is clear, with no evidence of cloudiness.
- Add 3 mEq/L of K if serum K level is 4 mEq/L.

G. Monitoring

- Maintain hourly PD flow sheet.
 - \circ Volume in
 - \circ Volume out
 - Net/hr (+/-)
 - Net over the course of dialysis (+/-)
 - Intakes (enteral, parenteral)
 - Outputs (urine, gastric, insensible water loss, etc.)
- Establish a desired fluid balance. Proceed gently if negative balance is required. Reassess the state of hydration frequently.
- Measure serum glucose and potassium every 4 hours for the first 24 hours or until stable, then twice a day. Obtain other serum electrolyte levels twice daily. Check blood urea nitrogen, serum creatinine, serum calcium, serum phosphorus, and serum magnesium once a day.
- Obtain cell count, Gram stain, and culture of peritoneal effluent every 12 hours.
- Recognize that some drug dosages may need adjustments (18,19 and 20) (see Appendix E).

H. Complications See Table 50.1.

Continuous Arteriovenous Hemofiltration in Newborns

A short discussion of CAVH and CVVH is included for completeness. However, use of these modalities should be limited to regional centers and performed by those with the required expertise.

TABLE 50.1 Complications of Peritoneal Dialysis	
Problem (Risk)	What to Do
Perforation of bladder, bowel, or major vessels (3%–7%)	Surgical consultation
Puncture-site bleeding (3%–15%)	Apply pressure gently.
	Purse-string suture.
Blood-stained dialysis maintained after several cycles	Check hematocrit frequently.
	Continue heparin.
	Rule out major-vessel bleeding.
Leakage from exit site (2%–20%)	Reduce dwell volume until leakage stops.
Extravasation of dialysate into the anterior abdominal wall	Replace with new catheter.
More than 10% of solution retained in each of several	Reposition infant gently.
consecutive cycles (outflow obstruction) (15%–30%)	Reposition catheter by rotation and slight
	retraction. Do not advance.
	Remove if unchanged.
	Replace with new catheter.
Two-way obstruction (3%–20%)	Irrigate catheter with small amount of dialysate
	or saline aseptically.
	Reposition.
	Remove if unchanged.
Dislodgment of catheter (3%)	Replace with new catheter.
Hydrothorax (0%–10%)	Reposition infant, head and chest above level of
	abdomen.
	Decrease dwell volume.
Hyperglycemia (10%–60%)	Avoid high concentrations of dialysate unless
	outflow is inadequate.
	Low dose of insulin if needed.
Lactic acidosis	Use bicarbonate dialysate ^a
Hyponatremia	Reduce fluid intake. Aim to increase outflow if
	secondary to fluid overload.
Hypernatremia	Increase fluid intake if secondary to excessive
	ultrafiltrate.
Exit site infection (4%–30%)	Systemic antibiotics.
Peritonitis (0.5%–30%)	Several rapid flushing exchanges.
	Blood culture. Systemic vancomycin plus
	ceftazidime or an aminoglycoside.
	For fungal peritonitis, systemic therapy is
	needed and catheter should be removed.
Hernia (inguinal or umbilical) (2%–13%)	Possible need for future repair.
Small bowel herniation and gangrene at catheter exit site (1	Surgical consultation
case report)	
Removal of therapeutic drugs	See Appendix E.
^a 1.5% bicarbonate dialysis solution: 140 mEq/L Na, 110 mE	Eq/L Cl, 30 mEq/L HCO3, 15 g of glucose; add
sterile water to 1,000 mL.	

Data from refs. (4,15,16,21,22,23,24,25,26,27,28,29,30,31).

CAVH is an extracorporeal technique for removing plasma water and its dissolved solutes of less than 50,000 Da over an extended period of time. With use of an arterial access line of the largest possible diameter and a venous access line, blood enters the extracorporeal circuit (arterial tubing, hemofilter, and venous tubing) by way of the arterial line and returns to the patient by way of the venous line (Fig. 50.4). The arteriovenous pressure gradient frequently produces adequate blood flow through the circuit; however, the addition of a blood pump may be necessary. As blood flows through the extracorporeal device, plasma water and dissolved solutes are filtered out (ultrafiltered) through the pores of a hemofilter. A hemofilter is composed of many fine capillaries of highly water-permeable membranes located within a cylindric case. The filtered-off fluid (ultrafiltrate) is drained out by way of an exit incorporated on the surface of the hemofilter. The fluid removed has all the characteristics of an ultrafiltrate of plasma water. Within the past few years, except when it is incorporated in an extracorporeal membrane oxygenation circuit for ultrafiltration, CAVH has been widely replaced by CVVH. Two single-lumen catheters (or one double-lumen venous catheter) are used for vascular access in CVVH. Blood flow is maintained by a pump and is therefore independent of the patient's systemic blood pressure. Several brands of CVVH machines are currently available in many pediatric centers for CVVH/D (32).



FIG. 50.4. A continuous arteriovenous hemofiltration circuit.

References

 Blatz S, Paes B, Steele B. Peritoneal dialysis in the neonate. Neonatal Network. 1990;8:41.
 Stapleton FB, Jones DP, Green RS. Acute renal failure in neonates: incidence, etiology and outcome. Pediatr Nephrol. 1987;1:1314. 3. Meeks ACG, Sims DG. Treatment of renal failure in neonates. Arch Dis Child. 1988;63:1372.

4. Matthews DE, West KW, Rescorla FJ, et al. Peritoneal dialysis in the first 60 days of life. J Pediatr Surg. 1990;25:110.

5. Fischbach M. Peritoneal dialysis prescription for neonates. Peritoneal Dial Int. 1996;16:S512.

6. Esperanca MJ, Collins DL. Peritoneal dialysis efficiency relation to body weight. J Pediatr Surg. 1966;1:162.

7. Chan KL, Ip P, Chiu CSW, Cheung Y. Peritoneal dialysis after surgery for congenital heart disease in infants and young children. Ann Thorac Surg. 2003;76:1443.

8. Anand SK. Acute renal failure in the neonate. Pediatr Clin North Am. 1982;29:791.

9. Moghal NE, Embleton ND. Management of acute renal failure in the newborn. Semin Fetal Neonatal Med. 2006;11:207.

10. Batshaw ML, Brusilow SW. Treatment of hyperammonemic coma caused by inborn errors of urea synthesis. J Pediatr. 1980;97:893.

11. Gartner L, Leupold D, Pohlandt F, et al. Peritoneal dialysis in the treatment of metabolic crises caused by inherited disorders of organic and amino acid metabolism. Acta Paediatr Scand. 1989;78:706.

12. Daschner M, Schaefer F. Emergency dialysis in neonatal metabolic crises. Adv Renal Replace Ther. 2002;9:63.

13. Mattoo TK, Ahmad GS. Peritoneal dialysis in neonates after major abdominal surgery. Am J Nephrol. 1994;14:6.

14. Chadha V, Warady BA, Blowey DL, et al. Tenckoff catheters prove superior to Cook catheters in pediatric acute peritoneal dialysis. Am J Kidney Dis. 2000;35:1111.

15. Kohli HS, Barkataky A, Kumar RSV, et al. Peritoneal dialysis for acute renal failure in infants: a comparison of three types of peritoneal access. Renal Fail. 1997;19:165.

16. Kohli HS, Bhalla D, Sud K, et al. Acute peritoneal dialysis in neonates: comparison of two types of peritoneal access. Pediatr Nephrol. 1999;13:241.

17. Ronnholm KAR, Holmberg C. Peritoneal dialysis in infants. Pediatr Nephrol. 2006;21:751.

18. Trompeter RS. A review of drug prescribing in children with end-stage renal failure. Pediatr Nephrol. 1987;1:183.

19. Aandgil A, Srivastava RN. Drug prescribing in children with renal failure. Indian Pediatr. 1989;26:693. 20. Bennett WM, Blyth WB. Use of drugs in patients with renal failure. In: Schrier RW, Gottschalk CW, eds. Disease of the Kidney. 4th ed. Boston: Little, Brown; 1988:3437.

21. Kohli HS, Arora P, Kher V, et al. Daily peritoneal dialysis using a surgically placed Tenckhoff catheter for acute renal failure in children. Renal Fail. 1995;17:51.

22. Bunchman T. Acute peritoneal dialysis access in infant renal failure. Peritoneal Dial Int. 1996;16:S509.

23. Walle JV, Raes A, Castillo D, et al. New perspectives for PD in acute renal failure related to new catheter techniques and introduction of APD. Adv Peritoneal Dial. 1997;13:190.

24. Blowey DL, McFarland K, Alon U, et al. Peritoneal dialysis in the neonatal period: outcome data. J Perinatol. 1993;13:59.

25. Sizun J, Giroux JD, Rubio S, et al. Peritoneal dialysis in the very-low-birth-weight neonate (less than 1000g). Acta Paediatr. 1993;82:488.

26. Werner HA, Wensley DF, Lirenman DS, et al. Peritoneal dialysis in children after cardiopulmonary bypass. J Thorac Cardiovasc Surg. 1997;113:64.

27. Dittrich S, Dahnert I, Vogel M, et al. Peritoneal dialysis after infant open heart surgery: observations in 27 patients. Ann Thorac Surg. 1999;68:160.

28. Sorof JM, Stromberg D, Brewer ED, et al. Early initiation of peritoneal dialysis after surgical repair of congenital heart disease. Pediatr Nephrol. 1999;13:641.

29. Reznik VM, Griswold WR, Peterson BM, et al. Peritoneal dialysis for acute renal failure in children. Pediatr Nephrol. 1991;5:715.

30. Huber R, Fuchshuber A, Huber P. Acute peritoneal dialysis in preterm newborns and small infants: surgical management. J Pediatr Surg. 1994;29:400.

31. Wong KKY, Lan LCL, Lin SCL, Tam PKH. Small bowel herniation and gangrene from peritoneal dialysis catheter exit site. Pediatr Nephrol. 2003;18:301.

32. Menster M, Bunchman TE. Nephrology in pediatric intensive care unit. Semin Nephrol. 1998;18:330.

51-Neonatal Hearing Screening

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A. Definitions

- Otoacoustic emissions (OAE): A screening tool that measures sounds generated by a functioning cochlea. A probe containing a microphone delivers a sound stimulus into the ear and records the response that travels back out of the cochlea, through the middle ear, and into the ear canal.
- Automated auditory brainstem response (AABR): A screening tool that records auditory brainstem responses and compares them to a template representing typical results in neonates. Click sounds are delivered by earphones into the ear canal, and electrodes placed on the head and nape of neck register the auditory brainstem response. In addition to assessing middle ear and cochlear activity, this test evaluates the function of the auditory nerve and auditory brainstem.
- Auditory brainstem responses (ABR); may also be referred to as brainstem auditory evoked response (BAER): A diagnostic test used to predict type and severity of hearing loss. ABR testing is conducted after a failed screening measurement. Auditory brainstem responses are determined in each ear for both click and tone stimuli. These sounds are presented by air (earphone) as well as bone conduction. Severity of hearing loss is expressed in decibels and described as conductive, sensorineural, or mixed.

B. Purpose

- Identifying and providing early intervention for children with congenital hearing loss to minimize delay in speech and language development.
- Supporting accurate reporting by state of incidence of congenital hearing loss.

C. Background

- The incidence of congenital hearing loss in the general population is approximately 2 to 4 in 1,000 live births in the United States (1).
- The risk for hearing loss can increase substantially when infants are exposed to known perinatal risk factors for hearing loss, such as TORCH infections, significant hyperbilirubinemia, and ototoxic medications. As technology improves and the number of at-risk infants graduating from neonatal intensive care nurseries rises (2), the need for early identification of hearing loss and intervention becomes even more critical.
- Any infant who fails hearing screening needs to be immediately referred to a pediatric audiologist. Delaying diagnosis of hearing loss can lead to significant problems in language and speech acquisition. The Centers for Disease Control Early Hearing Detection and Intervention Program recommends that infants identified by a failed hearing screen be referred for a comprehensive audiology evaluation as soon as possible, and always before 3 months of age.

D. Indications

• Ideally, every newborn should receive a hearing screen before discharge from the hospital. In 1993, the National Institutes of Health (NIH) Consensus on Early Identification of Hearing Loss concluded

that all infants should be screened for hearing impairment prior to hospital discharge, if possible (3). The American Academy of Pediatrics officially endorsed these recommendations in 1999.

Over the past 13 years, widespread support has led to state-mandated or state volunteer programs throughout the United States. Currently, 39 states and the District of Columbia have passed legislation mandating universal newborn hearing screeningâ€"that is, hearing screening for every infant, regardless of background and risk factors (4).

• Infants who meet high-risk criteria for acquiring hearing loss warrant immediate hearing screening, as well as ongoing monitoring. Table 51.1 lists factors known to contribute to neonatal hearing loss (5). It is critical that infants with any of these risk factors be referred to audiology after discharge, even if they have passed their initial hearing screen, in order that they may continue to be monitored during the years of initial language acquisition and development.

TABLE 51.1 High-Risk Registry to Screen Selectively for Hearing Loss

- Illness or condition that requires admission of 48 hours or longer to neonatal intensive care unit
- Stigmata or other findings associated with a syndrome known to include sensorineural or conductive hearing loss
- Family history of permanent childhood sensorineural hearing loss
- Craniofacial anomalies, including those with morphologic abnormalities of the pinna and ear canal
- In utero infection, such as cytomegalovirus, herpes, toxoplasmosis, or rubella
- Parental or caregiver concern regarding hearing, speech, language, and developmental delay
- Postnatal infections associated with sensorineural hearing loss, including bacterial meningitis
- Neonatal indicators, specifically hyperbilirubinemia at a serum level that requires exchange transfusion, persistent pulmonary hypertension of the newborn associated with mechanical ventilation, and conditions that require the use of extracorporeal membrane oxygenation
- Syndromes associated with progressive hearing loss, such as neurofibromatosis, osteopetrosis, and Usher syndrome
- Neurodegenerative disorders, such as Hunter syndrome, or sensory motor neuropathies, such as Friedreich ataxia and Charcot-Marie-Tooth syndrome
- Head trauma
- Recurrent or persistent otitis media with effusion for at least 3 months

E. Contraindications

- Patient has significantly atretic or total lack of external ear canal: Refer directly to pediatric audiologist.
- Although it is certainly fair to rescreen an infant who has potentially failed screening because of excessive background noise, vernix in ear canal, etc., multiple rescreening attempts in hopes of eventually obtaining a "pass†are not recommended and can contribute to delayed identification of congenital hearing loss.

F. Limitations

- Infant hearing screening can be compromised by environmental noise (such as a busy intensive care unit) or infant movement. OAE screening, more so than AABR, is particularly affected by vernix occluding the ear canal, or middle ear pathology such as effusion (6).
- OAE screening, although less time-consuming to set up and conduct, has a higher "refer†(fail) rate than AABR. The refer rates for OAE screening alone have been cited in the literature as being between 5.8% and 6.5%, with refer rates using AABR screening around 3.2% (7,8). In particular, infants who are <48 hours old are more likely to have a "refer†result if screened with OAE, as the presence of vernix and debris in the ear canal can be a significant factor (9). There is no

standardization of newborn hearing screening protocols in the United States at this time, and both technologies are in wide use.

It is difficult to state accurately how many infants who "fail†their hearing screen actually do hear normally (have a "false-positive†result). This is due mainly to the fact that, for a variety of reasons, a significant number of patients who get a "refer†result do not follow up for diagnosis with an audiologist. The Joint Committee on Infant Hearing benchmark for diagnostic follow-up is 70%, and there are large programs that have shown steady improvement toward meeting or exceeding this standard (10,11). In sparsely populated areas of the country, where resources are limited, however, as many as 90% of infants requiring diagnostic assessment are lost to follow-up (12). Similarly, the number of infants who pass their hearing screen and ultimately end up with a diagnosis of hearing loss ("false-negative†result) is difficult to assess. Children whose hearing loss is not present at birth but develops later on ("late-onset†or "acquired†hearing loss) truly do pass their newborn hearing screenings. They are not "missed†by screening: Their hearing loss is different in nature than hearing loss that is present at birth (13).

G. Equipment

- An OAE screening system consists of a miniature microphone housed in a probe that emits either click or tone stimuli. This assembly is coupled to a computer for analysis of the sound in the ear canal and processing of the otoacoustic emission. Results may be automatically analyzed and interpreted as either "pass†or "refer†for each ear. Figure 51.1 shows an infant undergoing OAE screening.
- An AABR screening system consists of occlusive earphones that cover the ears and emit sound stimuli.

Electrodes placed on the head and the back of the neck detect electrical activity from the auditory nerve and brainstem. A computer registers samples of the electrical activity over a fixed period of time. The averaged responses are then compared to a normal newborn template to determine if the result is a $\hat{a} \in \hat{c}$ or a $\hat{a} \in \hat{c}$ for each ear. Figure 51.2 shows an infant undergoing AABR screening.



FIG. 51.1. An infant undergoing OAE screening.

H. Precautions

Care should be taken to attempt screening in a relatively quiet environment, as well as ensuring that the infant is resting comfortably and the ear canals are free from obvious debris.



FIG. 51.2. An infant undergoing AABR screening.

I. Special Circumstances

- Hearing parents whose infant does not pass a hearing screening: Parents are often quite concerned to learn that their infant has not passed a hearing screening. This can be especially stress-provoking for parents whose infant may have spent a good deal of time in a neonatal intensive care unit (NICU) and may be facing additional medical concerns upon discharge. It is extremely important to remember that a failed hearing screening is not a definitive diagnosis of hearing loss. It is an important indicator that the infant needs immediate referral to an audiologist for further detailed evaluation, which may or may not result in a formal diagnosis of hearing loss.
- Deaf parents whose infant does not pass a hearing screen: Deaf parents, especially culturally Deaf individuals who use American Sign Language and identify strongly with being members of the Deaf community, are often thrilled to find out that their infant may have hearing loss. This is a cultural identification: These parents are rejoicing in the fact that their infant is like them and will have a cultural place of significance in their social world. This is often in direct opposition to the traditional medical perspective on hearing loss (as a pathology that needs to be addressed and "fixedâ€), and the parental reaction can be frankly shocking for involved health care professionals. It is very important to realize that these infants of culturally Deaf parents are not facing the immediate crisis of delayed language development referred to earlier. American Sign Language is a well-researched, intact language (14,15) that is immediately accessible to an infant of Deaf parents. Although it is still extremely important to establish audiologic follow-up for these infants of Deaf parents who fail a hearing screen, it is also critical to respect the potential cultural implication for such families. These parents may be celebrating in a manner very similar to hearing parents who are happy that their infant has passed the hearing screening.

J. Techniques

Both OAE and AABR screening systems can be automated: An individual only needs to be appropriately trained to set up and apply the equipment. A computer processes the incoming information and gives a readout of the result, usually as pass or "refer.

K. Complications

OAE and AABR are considered to be noninvasive and safe procedures. Like any procedure that involves the application of electrode pads, mild superficial skin abrasions could possibly occur with the removal of the electrode pads after AABR testing (16).

References

1. Centers for Disease Control. Early Hearing Detection and Intervention Program. Available at: www.cdc.gov/ncbdd/ehdi/FAQ/questionsgeneralHL.htm#prev. Accessed January 21, 2006.

2. Kessenich M. Developmental outcomes of premature, low birth weight, and medically fragile infants. NBIN. 2003;3(3): 80–87.

3. Early Identification of Hearing Impairment in Infants and Young Children. NIH Consens Statement Online 1993 Mar 1–3 [cited January 21, 2006];11(1):1–24.

4. Status of State Universal Newborn and Infant Hearing Screening Legislation and Laws. Available at: www.asha.org/about/legislation-advocacy/state/issues/overview.htm. Accessed January 21, 2006.

5. Joint Committee on Infant Hearing. Year 2000 Position Statement: principles and guidelines for early hearing detection and intervention programs. Pediatrics. 2000;106:798–817.

6. De Michele A. Newborn hearing screening. eMedicine. 2005. Available at:

www.emedicine.com/ent/topic576.htm. Accessed February 13, 2006.

7. Vohr B, Oh W, Stewart E, et al. Comparison of costs and referral rates of 3 universal newborn hearing screening protocols. J Pediatr. 2001;139(2):238–244.

8. Clarke P, Iqbal M, Mitchell S. A comparison of transient-evoked otoacoustic emissions and automated auditory brainstem responses for pre-discharge neonatal hearing screening. Int J Audiol. 2003;42(8):443–447.

9. Lin H, Shu M, Lee K, et al. Comparison of hearing screening programs between one step with transient evoked otoacoustic emissions (TEOAE) and two steps with TEOAE and automated auditory brainstem response (AABR). Laryngoscope. 2005;115

10. Prieve B, Dalzell L, Berg A, et al. The New York State Universal Newborn Hearing Screening Demonstration Project: outpatient outcome measures. Ear Hearing. 2000;21:104–117.

11. Vohr BR, Carty LM, Moore PE, Letourneau K. The Rhode Island Hearing Assessment Program: experience with statewide hearing screening (1993–1996). J Pediatr. 1998:133(3):353–357.

12. Estimated Number of Infants Receiving Diagnostic Evaluation (2004). Available at:

www.cdc.gov/ncbddd/ehdi/2004/Eval_04_A_web.pdf. Accessed July 21, 2006.

13. Weichbold V, Nekahm-Heis D, Welzl-Mueller K. Ten year outcome of newborn hearing screening in Austria. Int J Pediatr Otorhinolaryngol. 2006;70:235–240.

14. Stokoe W. A Dictionary of American Sign Language on Linguistic Principles. Washington, DC: Gallaudet Press; 1965, rev. ed. 1976.

15. Stokoe W. Sign language structure. Annu Rev Anthropol. 1980;9:365–390.

16. National Library of Medicine, Health Services/Technology Assessment Text. Health Technology

Advisory Committeeâ€"Minnesota (Static collection) page. Available at: www.ncbi.nlm.

nih.gov/books/bv.fcgi?rid=hstat6.section.3839. Accessed January 21, 2006.

52-Management of Natal and Neonatal Teeth

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The occurrence of teeth in the oral cavity at birth or within the first 30 days of life is uncommon. Such teeth have been called natal and neonatal teeth, respectively. This distinction, however, is temporal and artificial. Relevant clinical inferences can be made by further describing these teeth as mature or immature based on the quality of dental tissue and degree of dental development (1). Hebling et al. classified natal teeth into four clinical categories (2) (Table 52.1).

The reported incidence of natal and neonatal teeth varies; a range of 1in 2,000 to 3,500 is widely accepted (3). However, in a study of 18,155 infants, the reported incidence of natal and neonatal teeth was 1:716 (4). About 85% of natal and neonatal teeth are mandibular incisors (5,6). However, there are case reports of natal teeth in the posterior regions of the alveolar process (3,7,8), thereby necessitating an examination of the

posterior region of the alveolar processes at birth for the presence of teeth. Further, 95% of natal and neonatal teeth are a member of the normal complement of the deciduous dentition (9); this implies that supernumerary natal and neonatal teeth are rare. Hence natal and neonatal teeth should usually be retained.

A. Etiology

- Superficial positioning of the primary tooth germ (10)
- Infection and malnutrition (10)
- Febrile illness (10)
- Maternal exposure to toxins (polychlorinated biphenols, polychlorinated dibenzofuran, polychlorinated dibenzo-p-dioxin) (11)
- Syndrome/medical condition (Table 52.2) (10)

B. Clinical Presentation (Figs. 52.1,52.2,52.3 and 52.4)

There is variability in the presentation of natal and neonatal teeth. Although some have normal crown shape and color, and are held firmly in the alveolar process, others present as discolored microdonts with hypermobility. The latter are the immature type of teeth. The management of the patient depends on the clinical presentation.

C. Clinical Assessment

Clinical assessment should include an assessment of the tooth, oral soft tissues, and the systemic disposition of the patient.

- Dental assessment
 - Mobility: Tooth mobility >1 mm is usually an indication for the extraction of the natal/neonatal tooth.
 - Color and shape of tooth: Discoloration and abnormal morphology indicate an immature natal/neonatal tooth, which usually will require removal.
 - Root formation: This can be assessed with a dental radiograph. However, a loose tooth is likely to be lacking in root structure, and is likely to exfoliate spontaneously and early, with the risk of aspiration.
- Soft tissue assessment
 - Ventral surface of the tongue: Riga-Fede disease, although not a disease, is the term given to an ulcerative granuloma formed on the ventral surface of the tongue. It results from irritation of the tongue by the sharp margins of the mandibular incisor.
 - Gingival tissue: Gingival tissue adjacent to the natal/neonatal tooth should be examined for presence of inflammation or granulomatous lesion, caused by irritation by the sharp cervical margins of an immature tooth.
- General assessment:

Table 52.2 lists the systemic conditions associated with higher incidence of natal/neonatal teeth. They should each be ruled out to ensure that a pre-existing medical condition is not overlooked.

D. Precautions

• Keep in mind that the initial question in management of natal teeth is whether extraction is indicated. Indiscriminate extraction of natal/neonatal teeth is discouraged (12).

TABLE 52.1 Hebling Classification of Natal Teeth

- 1. Shell-shaped crown poorly fixed to the alveolus by gingival tissue with absence of a root
- 2. Solid crown poorly fixed to the alveolus by gingival tissue with little or no root
- 3. Eruption of the incisal margin of the crown through the gingival tissue
- 4. Edema of gingival tissue with an unerupted but palpable tooth
- Natal and neonatal teeth should be differentiated from cystic lesions such as Bohn nodules and Epstein pearls, by palpation and location in the infant's mouth. Bohn nodules and Epstein pearls are firm and have a smooth, rounded surface. There will usually be several nodules/pearls, and they may be located on the posterior palate or mandibular ridge.



FIG. 52.1. Patient 1: Normal (edentulous) alveolar ridge in neonate.

- Prior to extraction, it must be confirmed that the patient has received the appropriate dose of vitamin K at birth (10). There has been one report of difficulty in achieving hemostasis by local pressure after the extraction of natal tooth. This patient received microfibrillar collagen hemostat over the extraction site (3). Current literature supports the extraction of the natal tooth at 10 days or later after birth, unless there is significant risk of aspiration (10).
- A detailed family history should be obtained, to rule out inherited coagulopathy.
- Following the extraction, the socket should be curetted to remove odontogenic tissue (see F, Complications).
- Long-term care: Whether the patient receives conservative restorative treatment or extraction, the parents should be encouraged to maintain regular dental appointments with a pediatric dentist. This enables monitoring of the extraction site, and parental guidance in oral hygiene practices for their infant.

TABLE 52.2 Conditions Associated with Higher Incidence of Natal/Neonatal Teeth

Ellis-Van Creveld syndrome Hallerman-Streiff syndrome Craniofacial synostosis Multiple steacystoma Congenital pachyonychia Sotos syndrome Cleft palate Pierre Robin anomalad

E. Technique

Nonextraction case

If the tooth is firm and appears of normal color and shape, extraction is not indicated.



FIG. 52.2. Patient 2: Hebling Classification #3 neonatal tooth; not indicated for extraction.



FIG. 52.3. Patient 3: Hebling classification #2 natal tooth; this tooth was extracted.

- Should the mother complain of discomfort while breast feeding, the use of a breast pump and bottling of milk should be encouraged.
- If the patient presents with Riga-Fede disease, a pediatric dentist should be consulted. The sharp margins of the tooth can be smoothed using photopolymerized dental composite restorative resin. This results in spontaneous resolution of the tongue lesion (13).
- Pain relief and faster healing may be accomplished by carefully applying Kenalog in Orabase (14).

Extraction Case

Extraction is indicated if there is hypermobility of the tooth, or if the tooth is of the immature type (malformed, discolored, lacking root development). These would be classified as class 1 or 2 by Hebling et al. (2).

- Equipment
 - 2 x 2-in gauze piece
 - Topical anesthetic
 - Blunt-nosed sterile surgical scissors
- Technique
 - Apply a tiny amount of topical anesthesia on the tissue attachment of the tooth.
 - \circ Hold the tooth with 2 x 2-in gauze square with your fingers and gently remove the tooth.

If the tooth has to be grasped with forceps, then the infant needs to be referred to a pediatric dentist for evaluation and possible extraction.

 \circ $\;$ Blunt-nosed scissors can be used to cut the tissue if it is too fibrous.



FIG. 52.4. Patient 3: The natal toothâ \in which was removed by grasping the tooth with gloved fingers holding the tooth with a 2 x 2-in gauze square.

- F. Complications of Extraction
 - Tissue tags comprising dental papilla and/or Hertwig's epithelial root sheath remain in the extraction socket (15). These tissues may continue to form dental hard tissues, that is, dentin and root structure (15). These aberrant dental hard tissues may interfere with the normal eruption of adjacent primary teeth (15).
 - The development of postextraction pyogenic granuloma (16) and hamartoma (17) have been reported.

References

- 1. Spouge JD, Feasby WH. Erupted teeth in the newborn. Oral Surg Oral Med Oral Pathol. 1966;22:198.
- 2. Hebling J, Zuanon ACC, Vianna DR. Dente natalâ€"a case of natal teeth. Odontol Clin. 1997;7:37.
- 3. Brandt SK, Shapiro SD, Kittle PE. Immature primary molars in the newborn. Pediatr Dent. 1983;5:210.
- 4. Kates GA, Needleman HL, Holmes LB. Natal and neonatal teeth: a clinical study. J Am Dent Assoc. 1984;109:441.

5. Zhu J, King D. Natal and neonatal teeth: a review of 24 cases reported in literature. J Pediatr. 1950; 36: 349.

- 6. Badenhoff J, Gorlin RJ. Natal and neonatal teeth. Pediatrics. 1963;32:1087.
- 7. Friend GW, Mincer HH, Carruth KR, Jones JE. Natal primary molar: case report. Pediatr Dent. 1991;13:173.
- Masatomi Y, Abe K, Ooshima T. Unusual multiple natal teeth: case report. Pediatr Dent. 1991;13:170.
 Howkins C. Congenital teeth. Br Dent Assoc. 1932;53:402.

10. Cunha RF, Boer FA, Torriani DD, Frossard WT. Natal and neonatal teeth: review of the literature. Pediatr Dent. 2001;23:158.

11. Alaluusua S, Kiviranta H, Leppaniemi A, et al. Natal and neonatal teeth in relation to environmental toxicants. Pediatr Res. 2002;52:652.

12. Watt J. Needless extractions. Br Dent J. 2004;197:170.

13. Slayton R. Treatment alternatives for sublingual traumatic ulceration (Riga-Fede disease). Pediatr Dent. 2000;22:413.

14. Seminario AL, Ivancakova R. Natal and neonatal teeth. Acta Medic (Hradec Kralove). 2004;47(4):229–233.

15. Nedley MP, Stanley RT, Cohen DM. Extraction of natal teeth can leave odontogenic remnants. Pediatr Dent. 1995;17: 457.

16. Muench MG, Layton S, Wright JM. Pyogenic granuloma associated with a natal tooth: case report. Pediatr Dent. 1992;14:265.

17. Oliveira LB, Tamay TK, Wanderley MT, et al. Gingival fibrous hamartoma associated with natal teeth. J Clin Pediatr Dent. 2005;29(3):249.

53-Relocation of a Dislocated Nasal Septum

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A. Indications

To avoid future surgery, breathing and feeding problems, epistaxis, malocclusion, and sinusitis (1) Fetal compression sufficient to cause some degree of nasal deformation is a frequent physical finding on early newborn examination and normally resolves within 48 hours of birth (Fig. 53.1). In some instances, intrauterine forces or pressure applied during delivery cause a true septal dislocation (Fig. 53.2)Fig. 53.2). The incidence of true septal dislocation ranges from 1% to 4% of births (1,2,3,4,5,6 and 7). The otolaryngology literature indicates that septal dislocation should be relocated within a few days of birth for the best outcome (1,3,8). To differentiate compression deformity from true septal dislocation, apply gentle pressure to the tip of the nose; a dislocated septum will move farther from the midline at the base, a compressed septum will not move from the midline at the base. A compressed nose can be restored to normal anatomy with gentle pressure; a nose with septal dislocation cannot.

B. Contraindications

- Presence of other nasal or midline congenital anomalies requiring more extensive treatment
- Posterior septal dislocation
- Nasal orifice too small to easily admit smallest septal forceps

C. Equipment

• Septal forcepsâ€"modified Walsham or other appropriately sized septal forceps (Fig. 53.3).

D. Precautions

- Reduction should be performed within the first 3 to 4 days after birth.
- Otolaryngology evaluation for refractory dislocations or associated facial abnormalities
- Adequate restraint of infant, especially the head
- Remember that many newborns are obligate nasal breathers; insertion of a large-bore nasogastric tube into the stomach or an oral airway, prior to the procedure, will serve to separate the tongue from the palate and to promote oral respiration.

E. Technique

- Place septal forceps into the nares on the anterior aspect of the cartilaginous septum, posterior to columella. Advance blades gently, approximately 0.5 to 1.0 cm. Do not advance past the inferior aspect of the middle turbinate; do not force (Fig. 53.4).
- Gently close the forceps onto the septum.
- Direct the pressure of the lower edges of the forceps blades toward the midline, to move the septum into alignment with the nasal groove on the vomer (spine)â€"a slight upward motion may be required to lift the inferior border of the septum over the side of the vomer into the spinal groove (can be compared to replacing a sliding door into the slider) (Fig. 53.4B, C).
- Re-examine to ensure adequate reduction.
- F. Complications
 - Hemorrhage
 - Damage to nasal structures, e.g., the turbinates, septum
 - Damage to skull baseâ€" resulting in cerebrospinal fluid (CSF) leak (if speculum inserted too far)
 - Persistent dislocation



A FIG. 53.1. Nasal compression without septal deviation. A: Shortly after birth, the nose is asymmetrical from simple compression with an angled septum at rest. B: The septum assumes its normal angle. (From Fletcher MA. Physical Diagnosis in Neonatology. Philadelphia: Lippincott-Raven; 1998:211.)



FIG. 53.2. A: At rest it is difficult to distinguish a true deviation. B: Attempts to restore normal anatomy are unsuccessful as the septum remains deviated at the base. (From Fletcher MA. Physical Diagnosis in Neonatology. Philadelphia: Lippincott-Raven; 1998:211.)

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FIG. 53.3. Walsham septal forceps.



FIG. 53.4. A: Landmarks of nasal anatomy. (From)Fletcher MA. Physical Diagnosis in Neonatology. Philadelphia: Lippincott-Raven; 1998:210.) B: Shows the cartilagenous nasal septum displaced to the left from the ridge on the vomer. Large arrows indicate the direction of turn of the forceps blades needed to replace the septum into the groove; small arrows indicate the concurrent upward pull. C: Shows the septum postreplacement.

References

1. Tasca I, Compradretti GC. Immediate correction of nasal septum dislocation in newborns. Am J Rhinol. 2004;18(1):47.

2. Podoshin L, Gertner R, Fradis M, Berger A. Incidence and treatment of deviation of the nasal septum in newborns. Ear Nose Throat J. 1991;70:485.

3. Silverman SH, Leibow SG. Dislocation of the triangular cartilage of the nasal septum. J Pediatr. 1975;87:456.

4. Jeppesen F, Windfield J. Dislocation of nasal septal cartilage in the newborn. Acta Obstet Gynecol Scand. 1972;51:5.

5. Bhattacharjee A, Uddin S, Purkaystha P. Deviated nasal septum in the newbornâ€"a 1 year study. Indian J Otolaryngol Head Neck Surg. 2005;57:304.

6. Gray LP. Septal and associated cranial birth deformities: types, incidence, and treatment. Med J Aust. 1974;1:557.

7. Kent SE, Reid AP, Brain DJ, Nairn ER. Neonatal septal deviations. J R Soc Med. 1988;81:132.

8. Pentz S, Pirsig W, Linders H. Longterm results of neonates with nasal deviation: a prospective study over 12 years. Int J Pediatr Otorhinolaryngol. 1994;28:183.

54-Lingual Frenotomy

Kathleen A. Marinelli

A. Definitions

• Lingual frenulum a fold of mucosa connecting the midline of the inferior surface of the tongue to the floor of the mouth (1)

Generally thin, membranous, and avascular in the newborn (Fig. 54.1)

- Ankyloglossia (tongue tie) a congenital oral abnormality, characterized by an abnormally short, thick, and/or tight lingual frenulum (1,2 and 3)
 - Ankyloglossia derives from the Greek agkylos crooked, and glossa tongue.
 - Many different variations of tongue tie
 - Differing degrees of severity
 - May restrict mobility of the tongue tip
- Lingual frenotomy (tongue clipping) a minor surgical procedure, appropriate for treatment of significant ankyloglossia in infants

Can be accomplished at the bedside or in an outpatient clinic setting (4,5 and 6)

- Frenuloplasty or frenectomy more complicated surgical procedures employing Z-plasty technique or with removal of tissue
 - Reserved for older children, adults, or infants with a complicated lingual frenulum, such as a thickened frenulum containing genioglossus muscle
 - Performed in the operating room, by a surgeon, under conscious sedation or general anesthesia

B. Purpose

- Lingual frenotomy performed when the presence of ankyloglossia restricts or impedes an infant's ability to suckle successfully
 - Most common in breastfeeding infants
 - Occasionally seen in infants using an artificial teat
- Other problems related to ankyloglossia that may manifest in older children and adults (2,3,7,8,9,10 and 11).
 - Mechanical problems

Gingival recession, mandibular diastema, malocclusion, prognathism, difficulty with intraoral toilet (licking the lips, sweeping food debris off teeth)

- Articulation errors in speech
- Social effects
 - Difficulty playing a wind instrument, licking an ice cream cone or lollipop, etc.
 - Can lead to social embarrassment and the need for the more complicated frenuloplasty procedure outside the neonatal period

C. Background

- There is much controversy surrounding ankyloglossia regarding:
 - Definitions
 - Range from vague descriptions of a tongue that functions with a less than normal range of activity to a specific description of a frenulum that is short, thick, muscular, or fibrotic (3) (Figs. 54.1 and 54.2)
 - A range of methods to describe and quantify tongue tie have been proposed, including methods of measuring the anatomic differences, to quantifying observations.
 - Clinical significance (1,2,6,12,13)
 - Prior to the introduction and widespread use of breast milk substitutes in the early twentieth century, breastfeeding was necessary for survival.
 - Release of tongue tie was commonly performed by the midwife at delivery (12,14).
 - Tongue tie does not generally pose a problem for the more passive process of bottle feeding.
 - With a decrease in breastfeeding rates, frenotomy became unnecessary.
 - With the current resurgence in breastfeeding, and increasing knowledge of the risks of breast milk substitutes, tongue tie is again emerging as an entity that interferes with successful breastfeeding.
 - A recent article surveying >1,500 pediatricians, otolaryngologists, lactation consultants, and speech pathologists concluded that there is little consensus among and within these groups regarding the significance or management of ankyloglossia (15).
 - Need for surgical intervention (2)
 - Some babies with tongue tie can breastfeed successfully with no surgical intervention (1,16).
 - Each breastfeeding dyad is a unique combination of many factors, including the infant's intraoral structures, adequacy of suckling, and the size, shape, and elasticity of maternal nipples.
 - An emerging body of literature suggests that for those mother–baby dyads who are experiencing difficulty breastfeeding associated with the presence of tongue tie,

frenotomy is a safe, effective, and immediate means of providing relief of symptoms and supporting breastfeeding (12,13,16,17,18,19 and 20).

- Timing of surgical intervention: To facilitate breastfeeding, it can be done in the first days of life, or anytime thereafter if problems emerge.
- Incidence of tongue tie ranges from 0.02% to 4.8% in various studies (1,2,11,15,17,20,21).
 - There appears to be a genetic predisposition in some families.
 - Most studies report approximately a 2:1 male predominance.
- In a recent study, 22% of 425 North American pediatricians surveyed indicated they had performed frenotomies; however, only 10% reported being taught the technique during residency (15).



FIG. 54.1. Newborn with significant ankyloglossia. Note heart-shaped tongue, inability to raise tongue tip toward roof of mouth.

D. Indications

- In the neonate, presence of ankyloglossia, usually in a breastfeeding infant, causing one or more of the following (11,16,17,21):
 - Maternal nipple trauma, pain
 - Poor latch
 - Ineffective suckling; continuous suckling
 - Weight loss or poor infant weight gain
 - Early weaning



FIG. 54.2. Grooved retractor used to raise tongue and visualize the frenulum. Notice how thin and membranous the anterior edge is.

E. Contraindications

• Presence of genioglossus muscle or vascular tissue in the frenulum with no thin membranous tissue for incision

Refer to appropriate surgeon for consideration for frenuloplasty.

F. Limitations

• If the difficulty with breastfeeding was not caused by the tongue tie, release of the tongue tie will not result in improvement.

Even when tongue tie is the cause, attention must be paid to latch and suckling after release to ensure the best outcome.

G. Equipment

- Blanket or towel for swaddling
- Sterile iris scissors
- Sterile grooved retractor (optionalâ€"see below) (Fig. 54.2)
- Gloves
- Gauze pads
- Topical anesthetic gel for oral use (optionalâ€"see below)
- Cotton swab

H. Precautions (Fig. 54.2)

- Ensure, by careful examination of the frenulum, that there is no vascular or muscular tissue in the field of incision. Transillumination may be used to enhance visualization.
- Avoid submandibular duct orifices lateral to the frenulum.
- Avoid the thicker, more posterior part of the frenulum, which carries the blood supply.

I. Technique (2,3,7,12,15,20,21) (Figs. 54.2 and 54.3)

- Place the infant on a firm surface.
- Firmly swaddle the infant in a blanket or towel with the arms swaddled in at the sides.
- Have an assistant standing at the head of the infant to stabilize the shoulders with their fingers while steadying the head with their palms.
- Stand on right side of infant if right-handed.
- Visualize the frenulum by positioning light source to the left of the infant, allowing essentially transillumination of the frenulum.
- Place two gloved fingers of the left hand below the tongue, on either side of the midline, or position a grooved retractor, to push the tongue up toward the roof of the mouth, exposing the frenulum.
- Inspect the frenulum for any vascular or muscular structures.

Frenotomy should be done only if the frenulum is thin, transparent, and with no other structures present.

- Utilization of local anesthesia (optional):
 - With no anesthesia, there is minimal, brief discomfort (6,11,12,15,20,22) because the frenulum is poorly innervated.

Babies frequently squirm with positioning but usually do not cry during procedure.

• Alternatively, topical anesthetic gel can be applied to the frenulum with a cotton swab.



FIG. 54.3. Grooved retractor used to raise tongue. Iris scissors make incision.

- Divide the membranous frenulum with iris scissors (Fig. 54.3):
 - Begin at the free border, proceed posteriorly, closer to the tongue than to the floor of the mouth.
 - Occasionally, a single cut will free the tongue sufficiently.
 - \circ Usually, 2 to 3 small, sequential cuts (1 to 3 mm) are required.

Each subsequent cut allows improved retraction and visualization for the next cut.

• Divide frenulum anterior to the vascular bundle until tongue is freed and can extend past lower alveolar ridge and lips (Fig. 54.4).



FIG. 54.4. After incision, minimal blood noted. Tongue now extends past lower alveolar ridge.

- Control any bleeding (usually minimal) with direct pressure applied with a sterile gauze pad.
- Inform mother that breastfeeding may resume immediately.

Mothers frequently note an immediate and dramatic improvement in breastfeeding, with reduced discomfort, improved latch, stronger suckling, and absence of the clicking sounds frequently produced by the tongue-tied infant when attempting to breastfeed.

- Antibiotic therapy is not required.
- Postoperatively, a white fibrin clot may form.

Reassure parents that this is not a sign of infection.

• Arrange follow-up in 1 to 2 weeks to check healing of the incision.

J. Complications (2,4,11,12,15,20,22)

- Extremely rare when performed by practitioner familiar and comfortable with the procedure
 - Excessive bleeding virtually never occurs unless deep lingual arteries and/or veins are severed.
 - Infection
 - Recurrent ankyloglossia due to excessive scarring
 - Generally less severe than original presentation
 - Often amenable to revision surgery
 - Glossoptosis (tongue swallowing) due to excessive tongue mobility

Theoretical concern; has never been reported in modern literature

References

1. Hall DMB, Renfrew MJ. Tongue tie. Arch Dis Child. 2005;90:1211.

2. Lalakea ML, Messner AH. Ankyloglossia: does it matter? Pediatr Clin North Am. 2003;50:381.

3. Kupietzky A, Botzer E. Ankyloglossia in the infant and young child: clinical suggestions for diagnosis and management. Pediatr Dent. 2005;27:40.

4. Hansen R, MacKinlay GA, Mansen WG. Ankyloglossia intervention in outpatients is safe: our experience [letter]. Arch Dis Child. 2006;91:541.

5. Naimer SA, Biton A, Vardy D, Zvulunov A. Office treatment of congenital ankyloglossia. Med Sci Monit. 2003;9:CR432.

6. Wallace H, Clarke S. Tongue tie division in infants with breastfeeding difficulties. Int J Pediatr Otorhinolaryngol. 2006;70: 1257.

7. Kummer AW. Ankyloglossia: to clip or not to clip? That's the question. ASHA Leader. 2005;10:6, 30.

8. Lalakea ML, Messner AH. Ankyloglossia: the adolescent and adult perspective. Otolaryngol Head Neck Surg. 2003;128:746.

9. Lalakea ML, Messner AH. The effect of ankyloglossia on speech in children. Otolaryngol Head Neck Surg. 2002;127: 539.

10. Marchesan IQ. Lingual frenulum: classification and speech interference. Int J Orofacial Myol. 2004;30:31.

11. Messner AH, Lalakea ML. Ankyloglossia: incidence and associated feeding difficulties. Arch Otolaryngol Head Neck Surg. 2000;126:36.

12. Griffiths DM. Do tongue ties affect breastfeeding? J Hum Lact. 2004;20:409.

13. Wright JE. Tongue-tie. J Paediatr Child Health. 1995;31:276.

14. Horton CE, Crawford HH, Adamson JE, et al. Tongue-tie. Cleft Palate J. 1969;6:8.

15. Messner AH, Lalakea ML. Ankyloglossia: controversies in management. Int J Pediatr Otorhinolaryngol. 2000;54:123.

16. Hogan M, Westcott C, Griffiths M. Randomized, controlled trial of division of tongue tie in infants with feeding problems. J Pediatr Child Health. 2005;41:246.

17. Ricke LA, Baker NJ, Madlon-Kay DJ, et al. Newborn tongue tie: prevalence and effect on breast-feeding. J Am Board Fam Pract. 2005;18:1.

18. Marmet C, Shell E, Marmet R. Neonatal frenotomy may be necessary to correct breastfeeding problems. J Hum Lact. 1990;6:117.

19. Notestine G. The importance of the identification of ankyloglossia (short lingual frenulum) as a cause of breastfeeding problems. J Hum Lact. 1990;6:113.

20. Masaitis NS, Kaempf JW. Developing a frenotomy policy at one medical center: a case study approach. J Hum Lact. 1996;12:229.

21. Ballard JL, Auer CE, Khoury JC. Ankyloglossia: assessment, incidence, and effect of frenuloplasty on the breast-feeding dyad. Pediatrics. 2002;110:e63.

22. Amir LH, James PJ, Beatty J. Review of tongue-tie release at a tertiary maternity hospital. J Pediatr Child Health. 2005;41:243.

Appendices

Appendix A: Chapter 5

TABLE A.1 Recommended Dosages and Oral/Parenteral Ratios for Opioids

Drugs	Routes of Administration	Parenteral Dosage (mg/kg)	Frequency	Oral/Parenteral Ratio	Frequency
Morphine	i.m., i.v., s.q., p.o., p.r.	0.05-0.2	q 1–2 h i.v. a 2–4 h i m /s a	3-6	q 4–6 h ^a a 8–12 h ^b
Meperidine	i.m., i.v., s.q., p.o.	0.5-1.5	q 1–2 h i.v. q 2–4 h i.m./s.q.	4	q 4–6 h
Codeine	i.m., i.v., s.q., p.o.	0.5-1.0	g 2–4 h	1.5-2	q 4–6 h
Hydromorphone	i.m., i.v., s.q., p.o., p.r.	0.02-0.04	q 2–4 h	2–4	q 4–6 h p.o. q 6–8 h p.o.
Methadone Fentanyl	i.m., i.v., s.q., p.o. i.m., i.v., t.m./t.d. ^c	0.05–0.2 0.001–0.005	q 12–24 h q 1–2 h	2	q 24 h —

i.m., intramuscular; i.v., intravenous; s.q., subcutaneous; p.o., by mouth; p.r., per rectum; t.m./t.d., transmucosal/transdermal; q, every. "Pertains to regular oral preparations (e.g., MSIR [Purdue Frederick, Norwalk, CT, USA], Roxanal [Roxane Laboratories, Columbus, OH, USA], tablets and oral solutions).

^bPertains to slow-release oral or rectal preparations (e.g., MS Contin [Purdue Frederick], Duramorph [Baxter Pharmaceutical Products, Liberty Corner, NJ, USA], Roxanal SR). ^ct.m./t.d. preparations have not been standardized for use in term or preterm neonates.

Source: Vitali SH, Camerota AJ, Arnold JH. Anaesthesia and analgesia in the neonate. In: MacDonald MG, ed. Neonatology: Pathophysiology and Management of the Newborn. 6th ed. Philadelphia: Lippincott Williams & Wilkins; 2005:1557–1571, with permission.

TABLE A.2 Recommended Dosages for Nonsteroidal Anti-inflammatory Drugs

Drug	Dosage (mg/kg)	Routes of Administration	Frequency
Acetaminophen	10-20	p.o., p.r.	q 4–6 h
Aspirin	10-15	p.o.	q 4 h
Choline/magnesium trisalicylate	10-15	p.o.	q 6–8 h
Ibuprofen	5-15	p.o., p.r.	q 6–8 h
Ketorolac tromethamine	0.3-0.6	p.o., i.m., i.v.	q 6–8 h
Naprosyn	5-7	p.o.	q 8–12 h
Tolectin	5-7	p.o.	q 8–12 h

p.o., by mouth; p.r., per rectum; i.m., intramuscular; i.v., intravenous; q, every.

Source: Arnold JH, Anand KJS. Anesthesia and analgesia. In: Avery GB, Fletcher MA, MacDonald MG, eds. Neonatology: Pathophysiology and Management of the Newborn. 4th ed. Philadelphia: JB Lippincott; 1994:1334–1345, with permission.
Appendix B Chapter 31

TABLE B.1 Basic, All-Purpose Instrument Tray^a

- 1. Sponge forceps (two to three)
 - a. Straight or curved
 - b. Serrated 7-in
- 2. Medicine glass (one)
 - For antiseptic solutions, saline irrigation, or local anesthetic solution
 - b. l oz
- 3. Surgical drapes (one to three)
 - a. Single-use: transparent, adhesive; or
 - b. Sterile towels
 - (1) Fenestrated (circumcision)
 - (2) Fan-folded towels
- 4. Knife handle (one) or disposable scalpel
- 5. Scissors (one to two)
 - a. Fine or delicate scissors
 - b. Straight or slightly curved
 - c. Sharp/sharp (one) and
 - d. Sharp/blunt (one)
 - e. 4-51/2 in
- 6. Dressing forceps without teeth (three to four)
 - a. Eye or delicate
 - b. Half-curved (two) and
 - c. Straight (one to two)
 - d. 5½ in
- 7. Tissue forceps with teeth, (one to two)
 - a. Thumb or iris
 - b. 5½ in
- 8. Ring-handled tissue forceps, (one)
 - a. Allis or Babcock design
 - b. 6 in
- 9. Hemostatic forceps, ring-handled (two to three)
 - a. Fine or mosquito
 - b. Straight and curved
 - c. 5 or 51/2 in
- 10. Vein retractor with small-width blades (one)
- 11. Probe with eye, 51/2 in (one)
- 12. Needle holder (one)

^aSuggested quantities are within parentheses. Descriptors are given for ordering but are subject to individual preference. Disposable, single-use, or plastic instruments from commercially prepared trays are suitable for many procedures.

TABLE B.2 Selected Sutures Appropriate for Common Neonatal Procedure:

Туре	Raw Material	Tissue Use	Advantages	Disadvantages
Absorbable Chromic gut	Mammalian collagen with chromium salt to delay absorption	Subcutaneous (use plain gut for mucosa)	Retain strength for up to 90 d Can be used in presence of infection Requires only square-knot tie	Moderate tissue reaction (3+) Tends to fray when tied Faster absorption the smaller the caliber
Vicryl or Dexon	Synthetic copolymers	Subcutaneous fascia	Mild tissue reaction (2+) Low infectivity rate For absorbable suture or ligature Maintain knots	Cannot be used for approximation under stress 60% strength at 2 wk Safety in cardiovascular tissue not established Requires flat and square ties with extra throws
Nonabsorbable Silk	Braided protein filament from silk worm	Skin fascia	Best knot holding Easiest to use Strong for size Gone after 1–2 yr	High infectivity rate High tissue reaction (4+)
Nylon	Polyamide polymer Mono- or braided filament	Monofilament: skin closure and plastic surgery Braided: any tissue	Inert Least tissue reaction (0–1+) Lowest infectivity Very strong for size	Poor knot holding, requires at least six ties Not as easy to handle
Polypropylene Prolene	Polymer of propylene Monofilament	Skin Pull-out subcuticular Low infectivity Very strong for size Holds knot better than nylon Easier to tie	Inert Low tissue reaction (0–1+)	Remains encapsulated Completely absorbable
Mersilene Dacron	Polyester polyethylene terephthalate	Skin	Easy to use Low infectivity rate Holds knot well Good overall structure	Mild reactivity (2+)
Skin closure tape	Reinforced nylon filaments to back or porous paper tape	Skin superficial laceration or when subcuticular suture also used	Easy to place and remove Quick to apply No skin reactivity Least scarring No anesthetic required	Will not stick to wet or oily skin (wipe skin with alcohol first) Will not hold if wound is widely separated or under tension Cannot evert wound edges

Appendix C: Chapter 41.

TABLE C.1 Blood Products

	Shelf Life	Advantages	Disadvantages	Comments
Product	Shelf Life	Advantages	Disadvantages	Comments
Whole Blood A. Whole blood Hct 40%	1. CPD = 21 d 2. Heparin = 24-48 h 3. CPDA-1 = 35 d	 Provides volume Provides RBCs Provides some ceagulation factors 	 WBC, and platelets relatively nonfunctional unless fresh and unrefrigerated Storage lesion defects (K⁺) in plasma fraction 	 Used for exchange transfusion PRBCs and FFP preferable for correction of massive blood loss Hepatized blood not licensed for use in USA, but used in other countries
B. Quad pack collection Het 40%	1. CPD = 21 d 2. CPDA-1 = 35 d	 Allows multiple transfusions to one infant Volume of each quad adjustable Can be made into PRBCs 	 Outdate rates high unless NICU has number of infants Some wastage expected 	 Many neonatal units find this collection system valuable. Potential for small- volume collection by decreasing antico- agulant volume with ratio of 14:100 of anticoagulant/blood
C. Reconstituted whole blood Het variable	24 h	 Allows preparation of whole blood from stored RBCs (packed or frozen) and FFP Allows preparation of Group O cells with low- titer A and B antibody plasma 	1. Time for preparation	 Use for exchange transfusion Hematocrit may be adjusted by formula Provides replacement equivalent of fresh whole blood Formula for reconsti- tution: volume of plasma to add = volume PRBCs × (Het PRBCs/Het desired - 1).
D. Autologous fetal blood	 Fresh heparinized <4 h CPD = 21 d CPDA = 35 d Additive = 42 d 	 Potential immediate availability in delivery room or from blood bank 	 Risk of bacterial contamination Difficult to obtain correct anticoagulant blood ratio Requires anticipatory preparation for best procedural centrol Complicated procedure to perform in DR and maintain steribly 	 Information on advantages of autologous blood is limited Properly prepared and tested banked blood a better choice if time permits Developing countries exploring use Competition for umbil- ical cord banking Consider delaying cord clamping as alternative

Product	Shelf Life		Advantages	Disadvantages	Comments
Red Cell Products A. PRBCs Het 70%–90% in CPDA-1 Het 50%–60% in additive solutions	25	1. CPDA-1 = 35 d 2. Additive solutions = 42 d	 Readily available Easy to prepare 	 Accen tuated storage lesion defects if unit is at er id of shelf life Less R BC mass/mL of tran sfused produ it, if PRBCs in add tive solutions are us id 	 Principal use for correction of anemia With sterile connecting device, can remove aliquots for transfusion If only additive solution products available, hard pack for massive transfusion or for exchange transfusion
B. Sedimented RBCs Het 65%	250	As above	Does not require centrifugation	 Het m ay not be as high a c desired Conta ins more plasm; i 	
C. Leukocyte-depleted RBCs	250-350	Depends on anticoagulant- preservative solution as above	 WBC count <1-5 × 10⁶/product Reduces risk of transmission of CMV 	 Presto age LD preferi ed over bedsid e filters Canne et LD a unit from s ickle-trait donor Leuko depletion failure s occur 	 Indicated for prevention of febrile transfusion reactions and for leukocyte alloimmunization in older children, but these phenomena are rare in neonates Not indicated for prevention of GVHD
D. Washed RBCs	200	24 h	 Removes 80% WBCs Removes platelets, K⁺, anticoagulant Het adjustable—less viscosity 	 Time equired for preparation Equip nent not abvays available Expire s 24 h after washin g as "open" system 	 Can wash portions of quad pack May be combined with FFP for exchange transfusions Major indications include patients with hyperkalemia and T activation
E. Irradiated PRBCs	250-350	Depends on anticoagulant preservative solution	 Abrogates GVHD in susceptible infants 	 Reduc time d 2. Storag 28 d p or by c riginal expirat ion date Equip abways 	 Irradiation before issue preferable to long-term refrigerator storage postirradiation
F. Frozen deglyc- erolized RBCs	200	Frozen = up to 10 yr After thawing/deglycing 24 h	 Maintenance of 2, 3-DPG and ATP Removes >50% WBCs 	 Highe other Equip always Expire washin restations nent not available s 24 h after g 	 RBCs frazen with glycerol, thawed, and deglycerolized by washing prior to transfusion, resus- pended in 0.9% NaCI Allows storage of rare types of blood

Product	Shelf Life		Advantages	Disa	dvantages	Comments
Platelet Products A. Platelet concentrate (random donor)	45-50	5 d, depending on type of bag used	Approximately 5.5 × 10 ¹⁰ platelets in 45–60 mL plasma	 Conta WBCs and pl Rh D i possibl 	ins some , few RBCs, isma mmenization e	 Use platelet administration set Use immediately on receipt from blood bank and never refrigerate May be leukocyte- reduced but need 2–3 unit pool for efficient LD Volume may be reduced by centrifugation for use when out of group or extreme volume restriction is necessary; changes expiration to 4 h
B. Single-donor pheresis	300	5 d	 >3 × 10¹¹ platelets in 500 mL of plasma Always LD with current collection equipment If used over 5 d, can reduce donor exposures. Repeated pheresis from some donor possible 	1. Large to be a aliquoi	rolume, needs plit or ed	 Allows selection of HLA and PLA1 or other antigen- specific compatible donors With SCD, can remove aliquots for transfusion
Plasma Products A. FFP	190-300	Frozen (-15°C) = 1 yr Thawed = 24 h	 Contains plasma proteins, ceagulation factors, anticoag- ulant proteins, complement, and albumin 	 20–45 time Not for expans fibrino replace 	min thawing r volume ion or gen rment	 Separated from WB within 6-8 hours collection Must be ABO- compatible Use blood filter Confused with F24, now more com- monly available
B. Cryoprecipitate	10-25	Frozen ($-15^{\circ}C)=1$ yr Thawed = 4 h	 Better source of fibrinogen and VWF than plasma products 	 Limite includ conger fibrino deficie 	d indications e F XIII and iital/acquired gen ncy	 Must be ABO- compatible Use blood filter Transfuse immediately after Ordered in units but small volume confusing
C. Albumin	5%	3 yr at room temperature	 Heat-treated to reduce risk of infectious diseases Requires no cross- match Increases plasma oncotic pressure 	 Expension Does n coaguil 	se iot provide ation factors	 Five micron filter required Na = 200 Osmolarity 300 MOs m/L

Product	Shelf Life		Advantages	Disa	dvantages	Comments
Plasma Products			2001001	0.0000000000		
D. Albumin	25% 3 yr a ter	at room l mperature 2	 Requires no cross- match Increases plasma oncotic pressure with low volume 	 Expen Does r coagul Can p pulmo and ca 	se iot provide ation factors roduce nary edema, rdiac failure	 Five micron filter required Na = 130–160 Osmolarity 1,500 MOs m/L
E. Plasma frozen within 24 h (F24)	180–300 1 уг ро	if frozen; 1–5 d l sttfhawing	 Contains plasma proteins, coagulation factors, anticoagulant proteins, comple- ment, and albumin 	 May b for F V VWF 7 Antic protein Not in volum fibrino replace 	e less effective 'III, FV, and 'eplacement oagulant concentration dicated for e expansion/ gen :ment	 Separated from WB and frozen with 24 h Most commonly available from blood suppliers
F. Single source plasma	180-300	1	 Contains plasma proteins, coagulation factors, anticoagulant proteins, comple- ment, and albumin 			 From single-donor plasmapheresis Can be aliquoted into small volumes and frozen for neonatal use
G. Recovered plasma	180-300	1	 Contains plasma proteins, coagulation factors, anticoagulant proteins, comple- ment, and allournin 	 May b for F V VWF 1 7 Antic protein 	e less effective 'III, FV, and 'eplacement cagulant is	 Plasma recovered from WB without specialized time limit. Quality of factors/anticoagulant proteins not well studied

Appendix D: Chapter 34. Techniques for Endotracheal Intubation Specific to Unique Patient Needs

Elective Change of Orotracheal Tube in Intubated Patient

This procedure allows continued ventilation through a pre-established airway whenever it is necessary to change an endotracheal (ET) tube or to place a nasotracheal tube. By maintaining the original airway as long as possible during the change, there is less need for haste and less stress to the patient. An obvious prerequisite is that the original ET tube be patent and correctly positioned in the trachea.

Rapid Replacement Method

- Prepare equipment and patient as for initial orotracheal intubation.
- Release tube fixation device without displacing tube.
- Have assistant hold first ET tube in place at far left of the infant's mouth while continuing to ventilate infant.
- Visualize glottis with laryngoscope.
- Pass second orotracheal tube down far right of the mouth until it aligns with glottic opening.
- When new tube is positioned for direct insertion, have assistant withdraw first tube carefully.
- Advance new tube into position.
- Verify position and secure tube as previously described.

Alternative Method: Insertion over a Feeding Tube

Because of the narrow diameter of ET tubes in small infants, feeding tubes narrow enough to fit inside the ET lumen are often too flexible to stay within the trachea as the tubes are being changed. Be prepared to intubate directly should the feeding tube dislodge.

- Prepare equipment and patient as for initial orotracheal intubation.
- Release tube fixation device without displacing tube.
- Select the largest feeding tube that will easily go through the current and new endotracheal tubes. Remove the flared end of feeding tube and the adaptor on the new tube.
- Remove adaptor of currently in-place ET tube.
- Quickly insert the feeding tube through the lumen to a depth not greater than the ET tube.
- While holding feeding tube in place, pull ET tube out of trachea and off feeding tube.
- Slide new ET tube over feeding tube into trachea.
- Replace tube adaptor.
- Verify position and secure tube as previously described.

Selective Left Endobronchial Intubation

The angles of the bronchi are such that more often than not a tube will seek the right mainstem bronchus. The exceptions will be conditions that push the left side down (left upper-lobe emphysema) or that pull the right side up (marked upper-lobe atelectasis or hypoplasia). Normally, successful right mainstem intubation simply requires a longer tube. Selective intubation of the left bronchus is a more difficult and dangerous procedure; therefore, following all precautions is especially important.

Place the ET tube under guidance by direct bronchoscopy or under fluoroscopy when these procedures are available without compromise to infants (1,2).

The following procedure is a simple, indirect method based on a modification that tends to make the ET tube bend toward the left when it meets resistance at the carina (3).

- Cut an elliptical hole through half the diameter of ET tube 1 cm in length and 0.5 cm above the tip of the oblique distal end.
- Perform an orotracheal intubation as above, keeping the cut hole directed toward the left lung.
- Turn infant's head toward the right (4).
- While auscultating the lung fields, advance the tube to 0.5 to 1 cm below the calculated depth of the carina or until differential breath sounds are heard.
- If breath sounds diminish on the left, withdraw the ET tube until they return.
- Take a chest radiograph to confirm left bronchial position.
- Fix tube securely.
- Reassess position frequently, as tube may dislodge from one mainstem into the other.
- Follow patient closely for particular complications of
 - Air leak of ventilated area
 - Stasis pneumonia of nonventilated area
 - o Dislodgement from left mainstem bronchus
 - Ventilatory insufficiency due to significant disease in the only lung being ventilated

Nonvisualized Oral Intubation

This technique has a higher risk of complications and is less often successful than when direct visualization is used. Reserve the blind oral intubation for true emergencies in small infants when there is equipment failure (e.g., laryngoscope light) and when ventilation by mask is contraindicated (e.g., thick meconium).

- Stand at infant's feet.
- Carefully slide first two fingers of gloved, left hand into back of oropharynx at the base of tongue, until reaching vallecula and epiglottis. Keep fingers in the center of the tongue.
- Using index finger, pull epiglottis forward.
- Keep infant's head in midline.
- With right hand, guide ET tube, without stylet, along left middle finger, which is held just above index finger.
- Advance tube carefully just beyond fingertips.
- Avoid pushing against any obstruction.
- If available, have assistant press gently on trachea in suprasternal notch and report when tube passes under finger.
- Verify position, and fix tube as previously described.

Blind Nasotracheal Intubation (5)

Blind nasotracheal intubation is often used in adults. Because a stiff tube is needed, the chance of perforation in infants is greater if a stylet is used. Although an intubation under direct visualization is preferred, the presence of severe microagnathia or oral masses makes this approach valuable. It is critical not to push against any resistance.

- Keep infant supine with neck flexed and shoulders supported by a small roll.
- Shape a stylet so the tip of the endotracheal tube will curve anteriorly at 90 degrees. Be certain the tip of the stylet stays above the end of the ET tube. Alternately, freeze an ET tube in this configuration and remove stylet just prior to insertion.
- Maintaining the curve in the tube anterior, insert the tube carefully through the nostril until its tip is in the oropharynx.
- Pull the jaw forward into a sniff position with the head midline and put slight external pressure over the cricoid cartilage.
- Advance the tube to a suitable depth unless there is any resistance.
- Remove stylet and verify presence of exhaled humidity and equal breath sounds.

Intubation in Severe Cleft Defects

There are several possible modifications for ET tubes that are useful for fixation or elective intubation when there is a large cleft palate. For emergency intubations, the following modification using a standard tongue blade is usually immediately available (6). For techniques or difficult intubation alternatives, see above (7).

- Open infant's mouth and lay sterile tongue blade flat across maxilla, with ends extending from corners mouth. Have assistant hold in place.
- Follow steps for routine intubation, using tongue blade for support of laryngoscope as necessary.
- After intubation, fix tube to padded tongue blade.
- Recognize that tongue thrust on tube in absence of a normal palate may lead to extubation even without visible external lengthening of tube.

Emergency Retrograde Intubation (8)

When facial anomalies preclude other routes, retrograde intubation using a modified Seldinger technique is possible. Because the cartilaginous support of the trachea is so poor, needle puncture is far more difficult in neonates.

Equipment

- Venous cannula with stylet, 14 or 16 gauge
 - Feeding catheter. Verify that the catheter will pass through the lumen of the angiocath.
 - A 14-gauge cannula will admit a 5-French (Fr) feeding tube.
 - A 16-gauge cannula will admit a 3.5-Fr feeding tube.
- Hemostat
- Endotracheal tube

Technique

- Sedate infant if possible.
- Clean skin over cricothyroid area
- At the level of the cricothyroid, puncture skin with cannula and stylet. Angle cannula at 45 degrees from the skin and directed toward the head.
- Insert into lumen or trachea only until there is a give in resistance or air returns.
- Remove the stylet.
- Thread feeding tube through the lumen of the cannula until it can be retrieved from the nose or oropharynx.
- Bring cephalic end of feeding tube out of the nose or mouth, leaving other end well outside skin insertion.
- While feeding tube is in place, remove the cannula from the tracheal insertion site.
- Clamp the feeding tube at its tracheal insertion so it will not be pulled into the trachea farther than desired.
- At the upper end, slip the ET tube over the feeding tube and along its course until it has passed the proper distance into the trachea. Stabilize the ET tube.
- Cut the feeding tube at its tracheal insertion.
- While keeping the ET tube in place, pull the feeding tube through the ET tube.
- Secure ET tube after verifying correct intratracheal position.

References

1. Georgeson K, Vain N. Intubation of the left main bronchus in the newborn infant: a new technique. J Pediatr. 1980;96:920.

2. Mathew O, Thach B. Selective bronchial obstruction for treatment of bullous interstitial emphysema. J Pediatr. 1980;96:475.

3. Weintraub Z, Oliven A, Weissman D, Sonis A. A new method for selective left main bronchus intubation in premature infants. J Pediatr Surg. 1990;25:604.

4. Sivasubramanian K. Technique of selective intubation of the left bronchus in newborn infants. J Pediatr. 1979;94:479.

5. Williamson R. Blind nasal intubation of an apneic neonate. Anesthesiology. 1988;69(4):633.

6. Zawistowska J, Menzel M, Wytyczak M. Difficulties and modifications of intubation technique in infants with labial, alveolar and palatal clefts. Anaesth Resusc Intens Ther. 1973;1:211.

7. Stool SE. Intubation techniques of the difficult airway. Pediatr Infect Dis J. 1988;7:154.

8. Cooper CM, Murray-Wilson A. Retrograde intubation. Management of a 4.8 kg, 5 month infant. Anaesthesia. 1987; 42:1197.

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Appendix E: Chapter 50 TABLE E.1 Drugs Requiring Adjustment in Severe Renal Failure

Drug	Method	Adjustment	Elimination by PD
Acetaminophen	i	q8h	No
Acyclovir	i	q48h	No
Allopurinol	i	q12–24h	
	d	50%	
Amikacin	i	q24h	Yes
	d	20%-30%	
Amlodipine		Unchanged	
Amoxicillin	i	q12–16h	No
Amphotericin B	i	q24–36h	No
Amphotericin B cholesteryl	Unchanged		
sulfate			
Amoxicillin-clavulanic acid	Ι	q12–24h	No
Ampicillin	i	q12–24h	No
Bumentanide		Unchanged	
Calcitonin		Unchanged	
Captopril	d	50%	
Carbamazepine	d	75%	

Drug	Method	Adjustment	Elimination by PD
Carbenicillin	i	q24–48h	No
Cefaclor	d	33%	Yes
Cefamandole	i	q8–12h	No
Cefazolin	i	q24–48h	No
Cefotaxime	i	q12-24h	No
Cefoxitin	i	q24–48h	No
Ceftazidime	i	q24–48h	No
Ceftriaxone		Unchanged	No
Cefuroxime	i	q48–72h	Yes
Cephalexin	i	q12–24h	No
Cephalothin	i	q8–12h	Yes
Chloral hydrate		Avoid	
Chloramphenicol		Unchanged	No
Cimetidine	d	50%	No
Clavulanic acid	d	50%-75%	No
Clindamycin		Unchanged	No
Dexamethasone		Unchanged	
Diazepam		Unchanged	
Diazoxide		Unchanged	Yes
Dicloxacillin		Unchanged	No
Digitoxin	d	50%-75%	No
Digoxin	i	q48h	No
	d	10%-25%	No

Drug	Method	Adjustment	Elimination by PD
Diphenhydramine	i	q9–12h	
Enoxaparin		Not approved	
Erythromycin		Unchanged	No
Ethambutol	i	q48h	Yes
Fentanyl		30%-50%	
Flucytosine	i	q24–48h	Yes
	d	20%-30%	
Fluconazole	d	25%	Yes
Furosemide		Unchanged	
Gentamicin	i	q24–48h	Yes
	d	20%-30%	
Heparin		Unchanged	
Hydrocortisone		Unchanged	
Hydralazine	i	q8–16h	No
Indomethacin		Unchanged	
Insulin (reg)	d	25%-50%	
Isoniazid		Unchanged	Yes
Kanamycin	i	q24h	Yes
	d	23%-30%	
Ketoconazole		Unchanged	No
Labetalol		Unchanged	
Lidocaine		Unchanged	

Drug	Method	Adjustment	Elimination by PD
Lorazepam	d	50%	
Meperidine	d	50%	No
Metoprolol		Unchanged	
Metronidazole	i	q12–24h	No
Morphine		To avoid	
Nafcillin		Unchanged	
Naloxone		Unchanged	No
Nicardipine	d, i	Titrate	?
Oxacillin		Unchanged	
Penicillin G	i	q12–16h	No
	d	25%-50%	
Pentobarbital		Unchanged	
Phenobarbital	i	12–16h	Yes
Phenytoin		Unchanged	
Piperacillin	i	q24h	No
Prednisone		Unchanged	
Propranolol		Unchanged	

Drug	Method	Adjustment	Elimination by PD
Propranolol		Unchanged	
Ranitidine	d	50%	No
Rifampin		Unchanged	No
Secobarbital		Unchanged	No
Sodium nitroprusside		Unchanged	
Theophylline		Unchanged	Yes
Thiazide		Avoid	
Ticarcillin	i	q24–48h	Yes
Tobramycin	i	q24h	Yes
	d	20%-30%	
Valproic acid		Unchanged	No
Vancomycin	i	q24h	No
Verapamil	d	50%-75%	

Renal failure alters the clearance of most drugs to a degree that is inversely proportional to the glomerular filtration rate. Drugs that are entirely cleared by the liver are administered without renal adjustment. Dose is adjusted by either administerin a percentage of normal dose (d) or increasing the interval (i) between the doses by hours (18–20). The normal loading dose can be administered for virtually all drugs. Unlike hemodialysis, peritoneal dialysis usually has no significant effect on the clearance of most drugs. However, a supplemental dose is sometimes required. Blood levels: of drug, if available, are the best guide. PD, peritoneal dialysis; q, every.