CANCER DRUG DISCOVERY AND DEVELOPMENT

Biomarkers in Breast Cancer

Molecular Diagnostics for Predicting and Monitoring Therapeutic Effect

Edited by

Giampietro Gasparini, мо Daniel F. Hayes, мо



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BIOMARKERS IN BREAST CANCER

Molecular Diagnostics for Predicting and Monitoring Therapeutic Effect

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PREFACE

Biomarkers in Breast Cancer: Molecular Diagnostics for Predicting and Monitoring Therapeutic Effect is an updated view of the prognostic and predictive biomarkers in breast cancer written by experts in this field. This book covers the major advancements in the application of novel sophisticated molecular methods as well as the state of the art of the conventional prognostic and predictive indicators.

The first three chapters by Simon, Sweep et al., and Kimel et al. highlight the relevance of appropriate and rigorous study design and guidelines for validation studies on new biomarkers, concerning the standardization with quality control of the assay(s) used for their determination, their clinical development, and the statistical approaches. Of particular importance is the suggested optimized protocol for the HER-2/*neu* FISH assay applied by the NSABP network (1).

Gene expression profiling by tissue microarray is treated in depth in the following two chapters by De Bortoli and Briglia and by Kim and Paik. Recent studies conducted using this methodology have clearly documented the heterogeneous nature of invasive breast cancer within the same pathologic stage and menopausal status. These methods have clearly provided powerful new tools for more accurate individual definition of prognosis. However, much more work remains to be done before standard pathology laboratories can use tissue microarrays to perform tumor marker studies for routine clinical use.

Several individual factors are accepted or appear to be promising for standard clinical care. Of course, the use of estrogen and progesterone receptors (ER, PgR) to predict benefit from endocrine therapy represents the gold standard of tumor markers, and should be tested on every breast cancer tissue (2). More recently, testing for HER-2 status has also become standard to help select whether a patient with metastatic breast cancer should receive trastuzumab (2). Other factors that are still controversial, but are considered standard by some guideline panels, include indicators of cell proliferation as well as the urokinase-type plasminogen activator (PAI)-1 system. These are the topics of the next four chapters written by authors who are among the pioneers and worldwide experts in translational research studies in the field.

The roles of the epidermal growth factor receptor (EGFR) pathway and altered p53 in breast cancer growth and progression and as possible prognostic and predictive indicators are outlined by Ciardiello et al. and by Kandioler and Jakesz, respectively. In addition to trastuzumab, several preclinical and clinical studies suggest that other agents that disrupt the signaling pathways generated by members of the EGFR family may be effective against breast cancer. The ability to identify predictive surrogate biomarkers of response is the key for the rational selection of the patients most likely to benefit from these anti-EGFR compounds.

Moreover, it seems likely that markers of EGFR activity might provide the opportunity for monitoring therapeutic efficacy. The recent demonstration that mutation of the phosphorylation site of the receptor may be predictive of gefinitib activity in patients with non-small-cell lung cancer may represent an important step toward the right strategy for the use of such a class of new anticancer agents (3,4).

Adjuvant systemic therapy has now been clearly shown to reduce the odds of recurrence and death (5,6). However, because of the inaccuracy of currently available prognostic and predictive factors, much of adjuvant systemic therapy, especially chemotherapy, is given inefficiently, either to patients whose cancer was never destined to recur or to those patients whose cancer will recur, but for whom the therapy will not be effective. Enormous work has been directed toward prognosis, with advancements in the field of molecular biology and in the detection of occult metastatic cells distant from the tumor. Hawes et al. and Braun et al. comprehensively cover the methodology, state of the art, pitfalls, and promises of detection of early tumor cell dissemination in breast cancer patients.

Although determination of tumor biology in tissue justifiably garners much interest, the ability to test and monitor for biological changes with a simple blood test is obviously appealing. Chapters 13 and 14 deal with the clinical significance of circulating HER-2/neu and vascular endothelial growth factor (VEGF), these being among the more promising therapeutic targets for approaches based on selective molecular-targeting agents.

We hope this text offers a critical view of the modern approach to the development of surrogate biomarkers of prognosis and responsiveness to selective treatments in breast cancer. We are confident that it will provide useful reading for investigators involved either in laboratory research or in clinical development of prognostic/predictive indicators and of novel molecular-targeted therapy.

Giampietro Gasparini, MD Daniel F. Hayes, MD

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CONTENTS

Preface
Contributorsix
Part I. General Topics
 Guidelines for the Design of Clinical Studies for the Development and Validation of Therapeutically Relevant Biomarkers and Biomarker-Based Classification Systems
 Analytical Aspects of Biomarker Immunoassays in Cancer Research
3 Tissue Microarrays: State of the Art, Pitfalls, and Promises
Part II. Assays for Gene Expression and Post-Translational Protein Abnormalities
 Gene Expression Profiling with DNA Microarrays: <i>Revolutionary Tools to Help Diagnosis, Prognosis,</i> <i>Treatment Guidance, and Drug Discovery</i>
Part III. Assays for DNA Abnormalities
 5 Assays for Gene Amplification: State of the Art, Pitfalls, and Promises
Part IV. Tissue Predictive Biomarkers
6 Cell Kinetics
7 Urokinase-Type Plasminogen Activator and PAI-1: Validated Prognostic Factors for Breast Cancer

8	Predictive Value of c-erb-B2 for Endocrine Therapy	
	and Chemotherapy in Breast Cancer	
9	<i>erb</i> -B2 as a Therapeutic Target	
10	The Epidermal Growth Factor Receptor Pathway as a Selective Molecular-Targeted Treatment in Human Breast Cancer	
	Fortunato Ciardiello, Teresa Troiani, and Giampaolo Tortora	
11	<i>p53</i> as a Prognostic and Predictive Indicator	
	Daniela Kanaloler ana Kalmuna Jakesz	
	Part V. Predictive Biomarkers in Body Fluids	
12	Occult Metastatic Cells in Breast Cancer Patients: State of the Art, Pitfalls, and Promises	
	-	
13	Circulating HER2/neu: Clinical Utility	
14	Circulating Vascular Endothelial Growth Factor: Methods, Prognostic Significance, and Potential Application for Antiangiogenic Therapy	
	Roberta Sarmiento, Roberta Franceschini, Sabrina Meo, Massimo Gion, Raffele Longo, and Giampietro Gasparini	
15	Detection of Early Tumor Dissemination in Patients with Breast Cancer	

Part VI. Future Directions

16	New Technologies/New Markers/New Challenges		
Daniel F. Hayes and Giampietro Gasparini			
Inde	Χ		

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I GENERAL TOPICS

1

Guidelines for the Design of Clinical Studies for the Development and Validation of Therapeutically Relevant Biomarkers and Biomarker-Based Classification Systems

Richard M. Simon, DSc

CONTENTS

INTRODUCTION PITFALLS IN DEVELOPMENTAL STUDIES STRUCTURED RETROSPECTIVE STUDIES VALIDATION STUDIES REFERENCES

SUMMARY

Standards for the development of therapeutically relevant biomarkers and biomarker-based classification systems are lacking. The literature of prognostic marker studies for breast cancer is inconsistent, and few such markers have been adopted for widespread use in clinical practice. This is problematic, as many patients are overtreated and many others are treated ineffectively. The deficiencies in clinical development of biomarkers may become more severe as DNA

Cancer Drug Discovery and Development: Biomarkers in Breast Cancer: Molecular Diagnostics for Predicting and Monitoring Therapeutic Effect Edited by: G. Gasparini and D. F. Hayes © Humana Press Inc., Totowa, NJ microarrays and proteomic technologies provide many new candidate markers and therapeutics become more molecularly targeted. In this chapter we address some common problems with developmental marker studies and provide recommendations for the design of clinical studies for the development and validation of robust, reproducible, and therapeutically relevant biomarkers and biomarker-based classification systems. The design of validation studies is addressed for (1) identifying node-negative breast cancer patients who do not require systemic chemotherapy; (2) identifying node-positive breast cancer patients who do not benefit from standard chemotherapy; and (3) identifying node-positive breast cancer patients who benefit from a new molecularly targeted therapeutic.

Key Words: Biomarkers; microarrays; classification systems; clinical trial design.

1. INTRODUCTION

Breast cancer is a heterogeneous set of diseases. Although substantial progress has been made in the treatment of breast cancer, many patients are overtreated and many undergo intensive chemotherapy with little apparent benefit. The literature on prognostic factors in breast cancer, although voluminous, is inconsistent (1). The process of how to develop biomarkers that are robust, reproducibly measured, and therapeutically effective has not been well established. Although many prognostic factors have been studied, treatment selection has remained based primarily on the traditional components of Tumor-Node-Metastasis (TNM) stage and hormone receptor levels. This discrepancy between an inconsistent research literature and clinical practice will become even more problematic as DNA microarray and proteomic technologies provide new markers and therapeutics become more molecularly targeted. The objectives of this chapter are to provide information that facilitates the development of biomarkers for selection of the best treatment for each patient. We use the term biomarker to include predictive classification systems based on protein or RNA transcript profiles measured using technology such as DNA microarrays.

2. PITFALLS IN DEVELOPMENTAL STUDIES

Most biomarkers are developed using archived tumor specimens, and many of the problems that exist in the marker literature derive from the retrospective nature of these studies. Clinical drug trials are generally prospective, with patient selection criteria, primary end point, hypotheses, and analysis plan specified in advance in a written protocol. The consumers of clinical trial reports have been educated to be skeptical of *data dredging* to find something "statistically significant" to report in clinical trials. They are skeptical of analyses with multiple end points or multiple subsets, knowing that the chances of erroneous conclusions increase rapidly once one leaves the context of a focused single-hypothesis clinical trial. Marker studies are generally performed with no written protocol, no eligibility criteria, no primary end point or hypotheses, and no defined analysis plan. The patient population is often very heterogeneous and represents individuals for whom archived specimens are available. The patients are often not treated in a single clinical trial and represent a mixture of stages. Consequently, the overall population often does not represent a therapeutically meaningful group and the biomarkers identified may be of prognostic relevance, but less likely to be of predictive relevance for selecting therapy. Often the marker may be prognostic because it is correlated with disease stage or some other known prognostic marker. Broad populations are also often heterogeneously treated and so finding that a marker is prognostic in such a population may be difficult to interpret. Prognostic markers that do not have therapeutic implications are rarely used. The heterogeneous nature of the population also often results in multiple subset analyses of more therapeutically meaningful subpopulations. With multiple analyses, the chance of false-positive conclusions increases. Many biomarker studies perform analyses for many candidate biomarkers and several end points as well as for various patient subsets. Consequently, the chance for erroneous conclusions increases multiplicatively. The multiplicity problem is even more severe when one considers that there are usually multiple ways of quantifying biomarker level and many possible mathematical models for combining biomarker measurements.

Many of the problems that have hindered the development and acceptance of predictive single-protein biomarkers also apply for biomarkers based on DNA microarray expression profiles (2). There are multiple platforms and protocols for measuring expression profiles, and microarray research studies almost never evaluate interlaboratory assay reproducibility. Microarray expression profiles in research studies are generally performed at one time so that reagent variability is minimized, and it is almost never demonstrated that the models are predictive for tumor specimens collected and assayed at other times. This is of particular concern for printed cDNA microarrays where there may be substantial variability among batches of printed slides and batches of reference RNA.

Because of the number of genes available for analysis, microarray data can be a veritable fountain of false findings unless appropriate statistical methods are utilized. For example, in comparing expression profiles of 10,000 genes for tumor specimens selected from patients who have responded to a specified treatment to those for nonresponders, the expected number of false-positive genes that are statistically significantly (p < 0.05)

differentially expressed between the two groups is 500. This is true regardless of whether the expression levels for different genes are correlated. Consequently, more stringent methods for assessing differential expression must be used. Some studies do not use statistical significance at all and just identify genes as differentially expressed based on fold-change statistics; that is, the ratio of the average expression level in responders to the average in nonresponders, ignoring variability entirely. Others base their analyses on visual inspection of graphical data displays. Such methods are clearly problematic.

The unstructured nature of retrospective studies of biomarkers would not be so problematic if they were followed by structured prospective validation studies that tested specific hypotheses about predictive biomarkers. Such prospective trials are rarely performed, however, because they are difficult to accomplish. Consequently, before discussing the design of such prospective trials, we will make some suggestions about a more structured approach to retrospective studies.

3. STRUCTURED RETROSPECTIVE STUDIES

There is a role for exploratory studies in which multiple biomarkers and multiple ways of combining biomarkers into predictive models are examined so long as one has an adequate way of evaluating the result. A major problem with many retrospective studies is that they attempt to use the same set of data to both develop hypotheses (biomarkers) and to test those hypotheses. This problem is particularly severe when the number of candidate hypotheses examined in the exploratory stage is large.

In trying to determine which genes are differentially expressed in comparing responders to a given therapy to nonresponders, the number of hypotheses equals the number of genes examined. The Bonferonni method of adjusting for multiple testing requires that the p value calculated for comparing expression of a specific gene i in responders to nonresponders, say p_i , be adjusted based on the number of genes (N) examined. For microarray studies, N could be 10,000 or greater. The Bonferonni method tries to eliminate all false positives. For microarray studies, less conservative methods control the number of *false discoveries* (false positives), or the proportion of claimed positives that are false positives (*false discovery rate*) (3). These same ideas apply if, for example, we are examining which genes are prognostic for survival or disease-free survival on a particular treatment.

For assessing statistical significance, adjustments such as those described in the preceding paragraph can be applied to adjust for the fact that we do not have a specific hypothesis to test, but rather are in a hypothesis development mode. The adjustment is based on treating the problem as one of testing all possible hypotheses. For retrospective biomarker studies in which a number of biomarkers are examined, such adjustments to statistical significance should be applied. In many cases, however, statistical significance is not the best measure of biomarker value. A better measure is the extent to which the biomarker model enables us to predict whether the patient will respond to the treatment (4).

For binary outcomes such as response and nonresponse, the best measure of predictive accuracy is the number of correct predictions. For quantitative outcomes such as survival or disease-free survival, measurement of predictive accuracy is more complex. In many cases, it is reasonable to approximate quantitative outcomes in a binary manner: good outcome or poor outcome. In other cases, measures such as described by Korn and Simon (5) are used.

It is not valid to use the same set of data for selecting a predictive marker or developing a predictive model and for measuring predictive accuracy. The estimate of predictive accuracy computed on the same data used to select the marker or develop the model is called the *resubstitution* estimate and is known to be biased (6). The bias is extreme when the number of candidate markers is larger than the number of cases. For example, Simon et al. (6) showed that for two classes (e.g., responders and nonresponders) that have no genes that are truly differentially expressed in microarray expression profiles of thousands of genes, one can almost always find a predictive model that has a resubstitution estimate of accuracy of 100%. Such a model would be useless for future data, but would appear to give perfect predictions for the cases used to develop the model.

How can we develop a proper estimate of the accuracy of class prediction for future samples? For a future sample, we will apply a fully specified predictor developed using the data available today. If we are to emulate the future predictive setting in developing our estimate of predictive accuracy, we must set aside some of our samples and make them completely inaccessible until we have a fully specified predictor that has been developed from scratch without utilizing those set-aside samples.

To estimate properly the accuracy of a predictor for future samples, the current set of samples must be partitioned into a training set and a separate test set. The test set emulates the set of future samples for which class labels are to be predicted. Consequently the test samples cannot be used in any way for the development of the prediction model. This means that the test samples cannot be used for estimating the parameters of the model and they cannot be used for selecting the gene set to be used in the model. It is this latter point that is often overlooked.

The most straightforward method of estimating the accuracy of future prediction is the *split-sample* method of partitioning the set of samples into a training set and a test set as described in the previous paragraph. Rosenwald et al. (7) used this approach successfully in their international study of

prognostic prediction for large cell lymphoma. They used two thirds of their samples as a training set. Multiple kinds of predictors were studied on the training set. When the collaborators of that study agreed on a single fully specified prediction model, they accessed the test set for the first time. On the test set there was no adjustment of the model, redefining of cutoff values, or fitting of parameters. They merely used the samples in the test set to evaluate the predictions of the model that was completely specified using only the training data.

Cross-validation is an alternative to the split sample method of estimating prediction accuracy (8). Cross-validation can be used only when there is a well-defined algorithm for predictive model development. In such cases, cross-validation can be more efficient than the split-sample method for estimating prediction accuracy. There are several forms of cross-validation. Here we will describe leave-one-out cross-validation (LOOCV) in the context of a class predictor based on gene expression levels determined by DNA microarray analysis. LOOCV starts like split-sample cross-validation in forming a training set of samples and a test set. With LOOCV, however, the test set consists of only a single sample; the rest of the samples are placed in the training set. The sample in the test set is placed aside and not utilized at all in the development of the class prediction model. Using only the training set, the informative genes are selected and the parameters of the model are fit to the data. Let us call M₁ the model developed with sample 1 in the test set. When this model is fully developed, it is used to predict the class of sample 1. This prediction is made using the expression profile of sample 1, but obviously without using knowledge of the true class of sample 1. Symbolically, if \underline{x}_1 denotes the complete expression profile of sample 1, then we apply model M₁ to \underline{x}_1 to obtain a predicted class \hat{c}_1 . This predicted class is compared to the true class label c_1 of sample 1. If they disagree, then the prediction is in error. Then a new training set-test set partition is created. This time sample 2 is placed in the test set and all of the other samples, including sample 1, are placed in the training set. A new model is constructed from scratch using the samples in the new training set. Call this model M_2 . Model M_2 will generally not contain the same genes as model M_1 . Although the same algorithm for gene selection and parameter estimation is used, since model M₂ is constructed from scratch on the new training set, it will in general not contain exactly the same gene set as M₁. After creating M_2 , it is applied to the expression profile x_2 of the sample in the new test set to obtain a predicted class \hat{c}_2 . If this predicted class does not agree with the true class label c_2 of the second sample, then the prediction is in error.

The process described in the previous paragraph is repeated n times, where n is the number of biologically independent samples. Each time it is applied, a different sample is used to form the single-sample test set. During the steps, n different models are created and each one is used to predict the

class of the omitted sample. The number of prediction errors is totaled and reported as the leave-one-out cross-validated estimate of the prediction error.

At the end of the LOOCV procedure you have constructed n different models. They were constructed only in order to estimate the prediction error associated with the model constructed by applying the algorithm to the complete set of samples. The model that would be used for future predictions is one constructed using all n samples. That is the best model for future prediction and the one that should be reported in the publication. The cross-validated error rate is an estimate of the error rate to be expected in use of this model for future samples assuming that the relationship between class and expression profile is the same for future samples as for the currently available samples. With two classes, one can use a similar approach to obtain cross-validated estimates of the sensitivity, specificity.

Leave-one-out cross-validation is applicable only in settings in which there is an algorithm for the development of a predictive model. In many studies, the analysis is less algorithmic and many kinds of prediction models are explored. For such studies, it is best to use the split sample approach of setting aside at least one third of the samples as a test or validation set. The samples in the test set should not be used for any purpose other than testing the final model developed in the training set. Specifically, the test set samples should not be used for limiting the set of genes to be considered in detail in the training set. The samples in the test set should not be accessed until a single model is identified based on training set analyses as *the* model to be tested.

LOOCV can be used to evaluate risk group predictors using survival or disease-free survival data. Suppose we wish to identify patients in a low-risk group with 10-year disease-free survival >90%. Consider the leave-one-out training set in which observation *i* is left out in the test set. A disease-free survival model M_i is developed for the training set. For example, the model might be a proportional hazards regression model that predicts disease-free survival based on the expression profile and/or standard prognostic factors. The model M_i is applied to the left-out specimen *i* to obtain a prediction of the probability that the *i*th patient has 10-yr disease-free survival >90%. Let $y_i = 1$ if this probability is greater than 50%. This process is repeated for all of the leave-one-out training sets. Then, the Kaplan-Meier disease-free survival curve estimate is computed and plotted for the patients predicted to be of very low risk, those with $y_i = 1$. The adequacy of the model is judged by whether the estimated 10-yr disease-free survival for the identified lowrisk group is in fact in excess of 90%. An approach similar to this was used for developing a classification system based on survival for patients with renal cancer by Vasselli et al. (9).

One of the common errors in retrospective studies of biomarkers is that the statistical significance of the biomarker is evaluated rather than the predictive accuracy of the biomarker (4). We have indicated in the preceding how predictive accuracy can be evaluated in a manner that avoids the bias of the resubstitution estimate. But even this is not sufficient. New biomarkers are often correlated with existing prognostic factors. The retrospective study must provide strong evidence that the new marker is substantially more predictive than the currently available prognostic factors. This can be addressed by computing the split-sample or cross-validated error rate for a model consisting of current prognostic factors and then computing the split-sample or cross-validated error rate for a model consisting of current prognostic factors plus the new candidate markers. Only if the latter is substantially greater than the former with regard to a therapeutically relevant prediction will a prospective validation study be warranted.

4. VALIDATION STUDIES

Assuming that the initial study is performed properly with attention to the statistical principles described in previous section, it might be considered a phase II study, and the next step should be to conduct a phase III study that is focused on testing the specific classifier developed by the initial study (10). The phase III study should be conducted with a written protocol. The phase III trial should be designed to test the biomarker classifier developed in the previous study. The classifier should be fully specified in the protocol. If the biomarker is expression profile based, the specification must include the genes used, the mathematical form of the classifier, parameter values, and cutoff thresholds for distinguishing the classes or prognostic groups.

The phase III study should attempt to perform the assays in a manner as similar as possible to the way it would be performed broadly outside of a research setting if the diagnostic classifier were adopted. Consequently, attention is required in determining whether the same platform should be used for the phase III trial as for the phase II trial. If the platform is changed, then clearly some intermediate study will be needed to translate the classification algorithm from use on the phase II platform to the platform used in the phase III trial.

Even if there is not a change in platform, an intermediate study may be required to prepare the classifier for use with multiple laboratories performing the assay. In the phase II trial all of the assays may have been performed at a single location by a research laboratory and it may be advisable to conduct the phase III trial in a manner more similar to the way it would be performed if the classifier were adopted for national use. Generally this will mean that several laboratories will be conducting the assays. Consequently, the protocol for the phase III study should specify procedures to be used for conducting the assay. It is also useful to conduct intermediate studies of interlaboratory reproducibility of the assays. Unless interlaboratory repro-

Table 1 Guidelines for Validation Studies (10)

- 1. Intra- and interlaboratory reproducibility of assays should be documented.
- 2. Laboratory assays should be performed blinded to clinical data and outcome.
- 3. An inception cohort of patients should be assembled with <15% of patients nonevaluable owing to missing tissue or data. The referral pattern and eligibility criteria should be described.
- 4. Treatment should be standardized or randomized and accounted for in the analysis.
- 5. Hypotheses should be stated in advance, including specification of prognostic factors, coding of prognostic factors, end points, and subsets of patients and treatments.
- 6. The sample size and number of events should be sufficiently large that statistically reliable results are obtained. Statistical power calculations that incorporate the number of hypotheses to be tested and appropriate subsets for each hypothesis should be described. There should be at least 10 events per prognostic factor examined per subset analyzed.
- 7. Analyses should test whether new factors add predictiveness after adjustment for or within subsets determined by standard prognostic factors.
- 8. Analyses should be adjusted for the number of hypotheses to be tested.
- 9. Analyses should be based on prespecified cutoff values for prognostic factors or cutoffs should be avoided.

ducibility is sufficiently high, it is not advisable to proceed with the phase III trial.

If the biomarker classifier was developed using a dual-label microarray platform, then use of the classifier in other laboratories requires that they use the same common reference RNA as was used for the initial study. Because different batches of the common reference will be utilized for classifying subsequent patients, calibration studies will generally be required to ensure that the expression profile of the common reference does not change and to adjust the classifier for small changes.

Conducting the validation study as a prospective trial is desirable for many reasons. One can never be sure that the patients for whom one has adequate preserved tissue are representative of the population of patients presenting for treatment. It is difficult to assure that a retrospective cohort was adequately staged and treated, and the data available may be incomplete. It is also difficult to assess whether a diagnostic procedure is practical unless it is studied in the real-time context of presenting patients who need to be evaluated and treated. Prospective accrual is also important for evaluating the diagnostic classifier in the context of real-time tissue handling. Table 1, reprinted from Simon and Altman (10), indicates some important design features of prospective validation studies.

The objective of the validation trial of a predictive marker is to test the hypothesis that the marker is useful for treatment selection. This is often a more complex objective than validation of a prognostic marker, in which the objective is to determine whether the marker can separate the uniformly staged and treated patients into groups of differing outcome. There are some cases, discussed below, in which prognostic markers are also predictive markers. Our focus is on predictive markers and we will consider three breast cancer scenarios.

4.1. Identifying Node-Negative Patients Who Do Not Require Chemotherapy

Our first scenario is a putative marker for identifying node-negative patients whose prognosis on local therapy and possibly tamoxifen is so good that they do not require chemotherapy. The retrospective study for development of such a marker would have probably been based on archived tumors of node-negative patients who did not receive chemotherapy. A tissue microarray of a large number of such specimens, with associated clinical follow-up data, can provide a valuable resource for ensuring that the marker is sufficiently promising to warrant evaluation in a prospective clinical trial if the classifier is not RNA transcript profile based. A marker of this type meets the definition of a prognostic marker, but it can also be a predictive marker if it enables us to determine which node negative patients do not require chemotherapy.

The theoretically optimal trial design would be to randomize candidate node-negative patients to receive or not receive chemotherapy and then to validate whether the marker identifies those who do not benefit from chemotherapy. The candidate node-negative patients might be those with tumors 1–3 cm in diameter without known poor prognostic features such as hormone receptor negativity. This is probably not a feasible approach, however, because chemotherapy has already been established as being effective for much of the candidate population.

An alternative study design is to withhold chemotherapy from a subset of node-negative patients selected based on marker status to be of particularly low risk. If their outcomes were sufficiently good relative to some standard, then the marker would be accepted as useful. The standard might be based on outcomes for node-negative patients that are similar with regard to standard prognostic factors in other studies. It may also be useful to compare outcome for the selected patients (M⁺) to the outcome for the patients of the same series who did not have such predicted good prognosis (M⁻). The latter patients would have received chemotherapy, but their outcome even with chemotherapy may not be as good as that of the M⁺ patients without chemotherapy. If that is the case, then the value of the marker for withholding chemotherapy will have been demonstrated. An alternative approach would be to randomize patients selected as low risk based on marker status (M^+) to either receive or not receive chemotherapy. The marker would be validated if the randomized trial demonstrated that there was no clinically significant benefit of chemotherapy in the selected subset of patients. This would have to be a very large clinical trial, however. The benefit of chemotherapy would be expected to reduce the hazard or recurrence only by approx 25%, and with a very low event rate this is equivalent to a very small difference in absolute disease-free survival. The randomized trial asks a different question than the strategy described in the previous paragraph. The randomized trial asks whether there is a benefit of treatment. For patients with a very good prognosis, however, a statistically significant treatment effect may be of questionable clinical significance. Consequently, the randomized trial may not answer the most relevant question.

Gasparini et al. (11) describe guidelines for the adoption into clinical practice of new prognostic markers for use in treatment selection for patients with node-negative breast cancer.

4.2. Identifying Node-Positive Patients Who Do Not Benefit From a Chemotherapy Regimen

Consider now a putative marker that permits the identification of patients who do not benefit from a chemotherapy regimen that has been standard treatment. Let T denote the chemotherapy regimen, and let S denote local therapy or local therapy plus tamoxifen. The retrospective study used to develop the marker may have been based on tissue from a randomized trial of T vs S. In some cases the marker may be based on finding a signature of patients who do not respond to T in metastatic disease trials.

The ideal validation trial would probably be a randomized trial of S vs T for patients with node-positive breast cancer. One could analyze such a trial by seeing whether the benefit of T vs S depended on the marker level. Such a trial would generally be impractical, however, because T or some other kind of chemotherapy is standard treatment for node-positive patients. It might be possible, however, to randomize patients to receive or not receive one or more courses of T preoperatively, and to correlate marker result with biological response to T as assessed from the surgical specimen.

A second strategy would be to use chemotherapy T on all patients after measuring the marker. One could then determine prospectively whether the marker level correlates with outcome. This is a strategy analogous to that recommended in Subheading 4.1. Here, however, one is trying to determine whether the marker identifies a group of such poor disease-free survival on standard treatment T that the chemotherapy is judged nonworthwhile even in the absence of a control group not receiving chemotherapy. This strategy may be less satisfactory for judging poor prognosis in absolute terms than it was in Subheading 4.1. for judging good prognosis.

A third strategy would be to randomize the patients to marker based vs non-marker-based therapeutic management. The non-marker-based management would assign T to all patients. The marker-based management would assign T to all except those predicted based on the marker to be nonresponsive (M^-). One way of conducting such a trial is to measure the marker only for those patients assigned to marker-based management. The value of the marker is determined by comparing the outcome for the marker-based management arm to the outcome for the non-marker-based management arm. This is, however, a very inefficient trial design. Because most patients in both arms of the trial will be receiving the same treatment, the average treatment difference will be very small between the arms and a huge sample size will be required. The situation is even more problematic because it is a therapeutic equivalence trial in the sense that failure to find a statistically significant difference leads to the adoption of the new treatment approach, in this case marker-based treatment assignment.

A better design is to measure the marker on all patients, and then randomize them to marker-based treatment vs non-marker-based treatment. The evaluation of the marker can be performed by comparing outcomes for the M⁻ patients who received chemotherapy T on the non-marker-based arm but treatment S on the marker-based arm. This will require a much smaller sample size than the design described in the previous paragraph. This design is essentially equivalent to randomizing the M⁻ patients to T or S.

4.3. Identifying Node-Positive Patients Who Benefit From a Specific Regimen

Our third scenario is that we have a putative marker that identifies patients whose tumors are responsive to a new regimen E when the standard chemotherapeutic regimen is T. Many new therapeutics have defined molecular targets and are developed in conjunction with an assay that measures the expression of the target. The most adequate validation study is often a randomized clinical trial in which both marker-positive and marker-negative patients are randomized to either standard treatment T or T plus the new regimen E. The trial should be large enough so that the new regimen can be evaluated separately in the M^+ and M^- subsets. This requires about twice as many patients as if the regimen T+E were to be evaluated overall, without reference to the marker.

If the biological relationship between the marker and the therapeutic is sufficiently strong, it may be difficult to justify including marker negative patients in the study. A randomized study comparing T to T+E for M⁺ patients may be very efficient for demonstrating the effectiveness of the new treatment E, but it will not really constitute a validation of the marker.

The development of the therapeutic, supported by the marker assay, may, however, be more important than validation of the essentiality of the marker for selecting patients.

The least desirable alternative would be to randomize patients between T and T+E without measuring the marker. If the marker is important, then such a trial design may be very inefficient for evaluating the therapeutic E, and of course, it provides no information for validating the marker.

The scenario described here is also applicable to the development of treatment regimens in which the molecular target is not known or not known with certainty. Instead of using an assay based on the expression of the putative target, one may use a DNA microarray–based classifier developed in phase II trials of metastatic disease patients for distinguishing responders from nonresponders to the new regimen E. If tumor specimens are available from patients treated with the standard treatment T as well as those treated with the new treatment E, the classifier can be developed to identify patients who are predicted to be more responsive to the new treatment E but not to standard treatment T.

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Analytical Aspects of Biomarker Immunoassays in Cancer Research

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CONTENTS

2

INTRODUCTION IMMUNOASSAY DESIGNS VARIABILITY IN TEST RESULTS ASPECTS OF BIOMARKER ASSESSMENT QUALITY ASSURANCE CONCLUSIONS REFERENCES

SUMMARY

Many difficulties associated with immuno(metric) assay kits designed for quantification of a particular biomarker arise from their variation in specificity and binding affinity of the employed antibodies. Other important sources causing varying assay results are the use of different standard preparations in these kits and the nonuniform preanalytical specimen processing procedures employed, each of which should be subjected to standardization. To improve the performance and comparability of assays, continuous interlaboratory external quality control procedures are needed. Such quality assurance

Cancer Drug Discovery and Development: Biomarkers in Breast Cancer: Molecular Diagnostics for Predicting and Monitoring Therapeutic Effect Edited by: G. Gasparini and D. F. Hayes © Humana Press Inc., Totowa, NJ programs provide a forum for expert laboratory investigators to discuss technical details and to exchange laboratory issues and related practical information. This chapter addresses some of these issues and presents initial analytical validation procedures of newly developed biomarker assays, the validation of already established assay procedures for routine use on a day-to-day basis, and finally discusses some aspects on adequate (external) quality control proficiency testing.

Key Words: Biomarkers; immunoassay; cancer; tumor markers.

1. INTRODUCTION

In many cases progression of cancer growth is rather slow and often it may take years for a malignancy to manifest clinically. Because early cancer detection is required to significantly reduce cancer mortality, screening procedures are needed that are highly specific (i.e., providing almost a 100% proportion of negative test results for a tumour marker in nondiseased individuals) and sensitive enough to detect malignancies at an early stage of development. Thus, the screening procedure should give assay results above a defined cutoff value in a reasonable proportion of early stage diseased persons. As yet, there are no assay procedures available that meet such a specification, although there is a growing public interest in improving early cancer detection. Ideally, determination in biological specimens of cancerderived analytes for a particular type of cancer not only should provide valuable information for initial diagnosis, but also should have prognostic value to guide the choice of treatment, and such a test should provide a reflection of the tumor burden of the patient, being predictive for recurrent disease after initial treatment, and of help in monitoring the course of the disease throughout time of follow-up. Each of these properties should contribute to more effective treatment of an individual patient and thus provide indispensable information for improving the quality of life and outcome of the disease by increasing disease-free and overall survival. Despite extensive research efforts in the last decades and numerous papers dealing with development and clinical testing of potentially promising biochemical markers, no assays are as yet available that are sensitive enough to convincingly detect any of the major types of cancer at the most early stage. Although an impressive number of biochemical markers with the capacity to predict disease recurrence and/or early death have been introduced, a comprehensive understanding of the tumor biological processes involved is still lacking.

1.1. Guidelines for Evaluating Clinical Value of Biomarkers

At present, cancer diagnosis is based mainly on clinical symptoms and confirmed by histomorphological findings. Application of biochemical markers in this process may have additional value but still, depending on the marker test applied, the reliability criteria may become less important. In case of screening and diagnosis ("rule-in-disease") specificity is of utmost importance (to avoid false-positive assay results leading to unfavorable and unnecessary medical examination and treatment), although it should be realized that increasing specificity of a test goes at the cost of decreasing its sensitivity (that should remain high enough to detect early-stage diseased individuals). If the purpose of the test is disease monitoring to detect recurrence during follow-up, precision should be high; providing a prognosis for treatment, the test should put emphasis on specificity and accuracy. All of these criteria are not well established and should become standard criteria for evaluation of biomarker assays and their clinical application. In line with this, Hayes et al. (1) proposed certain criteria to standardize the available biomarker information for clinical use in a biomarker utility grading system.

Currently, many biochemical markers of potentially prognostic value are intensively tested in multicenter clinical trials. Only a few of these show a benefit for predicting prognosis of node-negative breast cancer patients. To conclude that these newly developed biochemical markers have independent prognostic value over already known factors, McGuire and Clark (2) some years ago proposed strict guidelines for evaluating newly developed prognostic markers, addressing the biological role of the new factor as well as the extent of the sample size, the risk of sample bias, the appropriate testing system, the establishment of cutoff values in a training data set, and confirmation of these observations in an independent validation data set.

2. IMMUNOASSAY DESIGNS

Immunochemical assay procedures can be classified according to the kind of analysis (qualitative, semiquantitative, or quantitative), type of assay format (manual or automated) and assay system (liquid phase, solid-liquid phase, [non-lequilibrium], making use of (radioisotopic or nonisotopic) labeled markers (to detect the antigen-antibody complex) or nonlabeled markers (in which the antigen-antibody complex is detected without labeled markers). The term immunoassay refers to competitive methods while immunometric assays refer to noncompetitive, sandwich-type assay formats. As early as 1969 the first generation of binding assays emerged with development of a radioimmunoassay (RIA) for quantification of insulin antibody formation. Later, the evolution of technical developments led to nonisotopic labels (enzyme-, fluorescence, time-resolved fluoro, (chemo-) luminescence immunoassays, etc.), monoclonal antibodies, phase matrices, and two-site immunometric sandwich-type assay formats employing two or more antibodies that will bind the analyte at repetitive or different epitope binding sites. The enzyme-linked immunosorbent assay (ELISA) format is

a commonly used type of two-site sandwich type assay in clinical routine work. The analyte is allowed to react noncompetitively with an excess of immobilized ("capture") antibody (coupled to a solid phase) and after addition and washing off the excess amount of sample specimen, an excess amount of marker labeled ("signal") antibody is added to bind to another epitope of the analyte. The sandwich thus formed is provided with marker label proportional to the amount of analyte present in the sample. These quantitative assay formats have several advantages: large numbers of specimens can be processed in parallel, providing reproducible results with reasonable precision, sensitivity, specificity, and accuracy. A major advantage of immunometric assay formats over semiguantitative or qualitative techniques is provided by the quantitative endpoint as measured against a defined standard, although there are limitations. Often, the analyte standard is not well defined, the assay procedure is not fully validated prior to use in patient studies, or the possibility to make comparisons to a reference method is lacking. Strict measures of quality assurance or good manufacturing practice protocols are needed to ensure proper assay performance before they should be applied.

Because early detection of small breast lesions is becoming common practice, there is an increasing demand to measure correctly biomarkers in smaller pieces of tumor specimens obtained through fine needle aspiration, core biopsies, or cryostat sections. This implies that there is an immediate need for more sensitive techniques than the standard immunoassays available to date. Alternative approaches to ELISA are proteomic methods such as MALDI and SELDI TOF mass spectrometry (MS), tandem MS, plasma resonance techniques, and antibody chip technologies. Of course the same rigorous principles of quality assurance should be applied for these new methodologies as for the more conventional immunoassays.

3. VARIABILITY IN TEST RESULTS

Assay results are often heterogeneous because of variations in specimen composition, tissue processing, design and specificity of the employed assay, as well as the statistics used for analyzing the collected data. In each of these stages, intrinsic differences in molecular forms (isoforms) of the biomarker present in the tumor tissue are augmented by external causes. The sampling procedure (e.g., fine-needle aspirate, core biopsy, or large biopsy obtained during surgery), the source of tissue (fresh or frozen), storage conditions (time, temperature, freeze–thawing cycles, etc.), and tissue processing (cytosol fraction, membrane extracts) may severely influence the final assay results (3). Likewise, this also holds true for the quantification of biological markers in serum or plasma (4).

Variable design of immuno(metric) assays results in the generation of different test results because different kits incorporate a broad spectrum of

antibodies, sometimes with different antibody specificities and/or affinities. Also, the use of different standards and reference materials provided with the kits are a source of variations in test results. Furthermore, different data reduction processes and statistical techniques are used to analyze tumor marker data and this may lead to a variety of conclusions regarding the clinical interpretation. The computational data processing of laboratory results must be appropriate, uniform, and evaluated extensively (5). McGuire and Clark stated that the design of confirmatory clinical studies should be identical to that of the definite study (2). It is of most importance to note that this also applies to all laboratory steps including tissue storage and processing, the analytical procedures, and the subsequent data processing.

The number and diversity of biomarkers for assessment of cancer prognosis is expanding rapidly, as is the variety of analytical formats and procedures used for quantification. A substantial proportion of assays is based on immunochemical principles and there is a widespread use of nonvalidated assay formats in clinical research settings. Because many assays are poorly standardized and (external) quality control is lacking in most cases, nonvalidated assay results without provided certified guidelines for interpretation become available at a too preliminary stage of assay development. Thus, biomarker testing procedures in laboratories participating in clinical trials should be standardized and externally quality assessed. This requires settlement of quality standards of all assay reagents included in assay kits, provision of guidelines for standardized assay protocols, standardized algorithms for calculation of assay results, and statistical procedures to allow unequivocal interpretation of clinical effect measures. Finally, to ascertain continuity of reliable biomarker data generation, there is a need for guidelines toward uniform internal and external quality assessment procedures. The next sections will discuss preanalytical, analytical, and postanalytical aspects of assay performance.

4. ASPECTS OF BIOMARKER ASSESSMENT

4.1. Preanalytical Criteria

4.1.1. SAMPLING BIAS OF TISSUE SPECIMENS AND TISSUE PROCESSING

Because many tumors are heterogeneous the size of a tumor tissue specimen is important to avoid sampling bias. This bias may lead to different assay results if different areas of a tumor are analyzed (different content of tumor cells, nonmalignant cells, extracellular matrix, fat, and necrotic spots). Thus, fine needle biopsy results may differ from those obtained from a tumor tissue biopsy specimen. Selection bias may occur if frozen tissue specimens from large tumor banks are used in retrospective studies as generally in tumor banks relatively larger samples of frozen tumor tissues are overrepresented (6).

The use of blood specimens requires standardization of blood collection conditions (fasting, fixed time of day, supine position), type of specimen (whole blood, serum, or plasma) and type of anticoagulant. Care should be taken to immediately transport tissue specimens or blood directly after surgery or blood collection to the laboratory in a standardized manner (time, temperature). Disintegration or extraction procedures of tissue samples should be performed according to the consensus protocols written by internationally acknowledged experts. Errors in this preanalytical phase of biomarker level quantification will affect the reliability of the final experimental data.

4.2. Analytical and Reliability Criteria

Prior to producing and subsequent reporting of test results it is the task of the laboratory to verify or establish performance specifications for each analytical procedure, irrespective whether the assay of interest has been developed in an academic institution or by a commercial company. In their instructions for use, kit manufacturers have often included disclaimers for misusing or overinterpreting the information included in their product information. It is common practice of diagnostic kit manufacturers to advice their clients that each laboratory should establish its own reference values in particular for specified populations or applications, irrespective of already available data provided by the manufacturer. The next sections deal with reliability criteria of analytic testing systems.

4.2.1. STANDARD CALIBRATION PREPARATIONS

Standards are used to prepare a standard dose response curve that relates the response reading as the independent variable to the quantity of the standard as the dependent variable. This allows calculation of the quantity of analyte from the response reading obtained for the unknown sample. It is not always possible to obtain sufficient quantities of a reasonably pure biomarker for characterization, which is the reason why in many cases arbitrary nonpurified or semipurified preparations of biomarkers are used to produce standard curves. Protein analytes may be present in different molecular forms ("isoforms") which may cause differences in affinity or other binding characteristics with antibodies. In case there are differences in affinity of the antibody for the calibrator standard and the analyte present in the unknown sample, different assay results will be obtained at different sample dilutions. For this reason we propose to analyze biological markers in at least two or three different dilutions to detect this phenomenon. This means that the suitability of a biomarker assay has to be validated for each biological specimen of concern because the procedure for the measurement of an analyte in tissue extracts is not always suitable for assaying the same analyte in plasma or serum. Stability of the standard can best be followed by longitudinal monitoring of the consecutively produced slopes of the

standard dose–response curves. Thus, assays should use well-defined, wellcharacterized standard calibrator material with known sequence and degree of purity. Also, different kit manufacturers should adhere to internationally accepted standards and preferably use identical standards in their diagnostic kits. An important source of providing biological reference materials to the scientific community covering many areas of clinical medicine is the WHO International Laboratory for Biological Standards (National Institute for Biological Standards and Control [NIBSC], Potters Bar, UK). Finally, as an example of advancements that contribute to standardization of widely used biomarker assays, we mention the introduction of an assay procedure for prostate-specific antigen (PSA) that determines several molecular forms of PSA on an equimolar basis, and the calibration of this assay with the Stanford 90:10 Reference Material, composed of 90% PSA-ACT and 10% free-PSA (7).

4.2.2. ACCURACY

Definition of accuracy of an assay by the International Federation of Clinical Chemistry (IFCC) is the agreement between the best estimate of a quantity and its true value. As this quantity has no numerical value, the term inaccuracy is used. Thus, inaccuracy is the difference between the mean of a set of replicate measurements and the true value. Although the concept is clear, it has realistic value for those analytes for which a reference method is available. As no such reference values are available or even feasible for many biomarkers, the concept of (in)accuracy has limited significance, emphasizing even more the necessity of standardization of assays.

4.2.2.1. Linearity

As outlined earlier, linearity of an assay in fact refers to identity between affinity of the antibody for the calibrator standard and the analyte present in the unknown sample. This should be the case and these tests of parallelism between standard and unknown analyte can be conducted by measuring samples at different sample dilutions and multiplying the amount of analyte measured with the dilution factor. Linearity studies are used to assess and establish the working range of an assay that is in between the lowest and highest concentration that can reliably be measured with that assay. This can easily be realized by mixing two different samples in several proportions (e.g., 1:3, 1:1, 3:1). See also the National Committee for Clinical Laboratory Standards (NCCLS) evaluation protocol (EP6).

4.2.2.2. Recovery

Recovery experiments are conducted to test whether the standard of the assay and analyte in the unknown sample behave chemically identical, or to exclude whether disturbing interactions of the analyte with the matrix or other compounds of the assay will lead to different assay results. Thus, in order to obtain insight into the identity of the analyte vs the standard, or to study matrix interactions with the standard, known amounts of standard are added to samples with an already known amount of endogenous biomarker and the recovery of the added amount is calculated.

4.2.2.3. High-Dose Hook Effect

This phenomenon is a source of error specifically occurring in double determinant one-step sandwich-type assays and comprises the saturation of capture and/or signal antibodies resulting from extremely high concentrations of biomarker analyte present in the incubation medium. This leads to a falsely low concentration calculated for the analyte. High-dose hook effects can be avoided by conducting a two-step assay protocol in which the immobilized capture antibody is incubated with an appropriately diluted unknown sample and excess of unbound analyte is washed off. The assay is completed by addition of signal antibody in the second incubation step. It is also advised to analyze samples at different dilutions to check whether the assay is vulnerable for the high-dose hook effect.

4.2.2.4. Interferences

Heterophilic antibodies are an often underestimated source of error in immunometric assays. In particular the treatment of patients with monoclonal mouse antibodies for immune-imaging and immune-targeting purposes has emerged occurrence of human antimouse antibodies (HAMAs), that is, the generation of human immune globulins G and M (IgG, IgM) in the blood of these patients. These antimouse IgG or IgM may also originate from other iatrogenic animal sources, all of these interfering to variable extents with the antibodies incorporated in biomarker sandwich-type assays. For a review of HAMA occurrence and it consequences for assay methodology see Kricka (8).

4.2.3. Specificity

In epidemiological terms specificity refers to the proportion of truenegative test results of a control population and in fact is similar to its definition in analytical terms where it is defined as (absence of) interference (cross-reaction) in an assay system of compounds more or less related to the analyte to be measured in that assay. Thus, specificity of immunoassays refers to the degree of interference by compounds that may resemble but differ from the analyte to be quantified. One established manner to express cross-reaction is comparison of the amount of analyte homologous for the assay with the amount of another compound tested for interference with the assay. This is performed at half the maximum response level (often referred to as $B/B_0 = 0.5$) of the linearized standard dose–response curve. The specificity of immunometric assays strongly depends on antibody characteristics because polyclonal or monoclonal antibodies or mixtures of both are applied in different testing kits. Specificity will be highest with monoclonal antibodies because these are directed against one epitope on the analyte molecule. Many tumor-associated antigens have epitopes also common to other proteins present in a variety of many other tissues. Because epitope mapping data of antibodies is not often documented, investigators have to check cross reactivity of a number of compounds related structurally or biologically to the assay's analyte.

4.2.4. SENSITIVITY

In epidemiological terms, sensitivity refers to the proportion of truepositive test results of a diseased population. Analytically, sensitivity may be defined as the limit of detection of the analyte in the assay, that is, the lowest concentration of analyte significantly different from zero, also called the analytical sensitivity. The limit of quantification at which a test can be reliably measured with a coefficient of variation of less than 20% is called the functional sensitivity. It is recommended to report clinical assay results not below the functional sensitivity limit that can easily be retrieved from the precision profile of an assay that is constructed by plotting the coefficients of variation of replicate measurements of all the samples assayed against the concentrations of the obtained results. Data on sensitivity should be provided by the kit manufacturer and checked by the investigator on first use of a kit. One of the goals of immunoassay methodology is to optimize continuously the lower detection limits of assays in order to settle clinically relevant cutoff points. Defining such low thresholds requires a high degree of reproducibility of assay results, that is, precision.

4.2.5. PRECISION

According to the IFCC, precision is defined by the agreement between replicate measurements. As is also the case with accuracy, precision has no numerical value, the reason why the use of imprecision is more practical, although not commonly used. The imprecision is the standard deviation or coefficient of variation of the results of a set of replicate measurements.

The precision of a biomarker determination varies depending on whether duplicate determinations are performed in one sample, different samples in the same batch, or in different batches, and so forth. Obviously, the estimate of the precision used to assess the validity of experimental results must be related to the assay conditions in the definite study. For instance, if the concentration of a biomarker in malignant vs nonmalignant tissue of the same patient is determined in one assay run, the statistical significance of relevant difference is referred to the intraassay precision of the method. On the other hand, when a marker is monitored over a long time of observation (follow-up), samples will be assayed in different batches of test kits and the interassay precision is the more relevant parameter. For validation of an assay, at least the intrasample, intraassay precision performance should be investigated. The precision profile is an ideal tool to assess this (see below). The NCCLS offers a practical evaluation protocol (EP5) for evaluating the precision performance of an assay.

4.2.6. MINIMAL CONSISTENCY CRITERIA

Apart from the aforementioned assay characteristics that should be assessed by the investigator once a new kit is introduced into the laboratory, assay performance may be hampered by day-to-day, performer-to-performer, and batch-to-batch variability. Run-to-run performance errors may be reduced by daily consistency testing of the calibration curve, the precision profile, and data on quality control specimens. Charting of standard dose–response curve characteristics comprises at least the calibrated slope; *y*-intercept, correlation coefficient, analyte concentration at 50% response (ED₅₀), and minimum detectable analyte concentration. The shape of this curve defines the quality of the performed assay and offers a basis for selection of the working range of the assay, while it also quite easily allows to detect unreliably scattering duplicates.

4.3. Postanalytical Criteria

Once an assay has been performed the results of unknown samples must be derived from the obtained response parameters by calculating the analyte concentrations from the standard dose–response curve. Numerous computerized algorithms are available, but irrespective of the choices made, it is highly advisable to use the same statistical approach to process assay data, especially if one participates in or conducts a multicenter study. Each laboratory should establish its own reference values to circumvent population sampling errors and biological variation.

5. QUALITY ASSURANCE (QA)

Defined protocols for (internal and external) quality control (QC) should be part of routine practice in the laboratory. QA not only comprises the analytical process as such (QC), but it also regards the total of the managerial, technical, and interpretative aspects and is intended to prevent, monitor, and correct mistakes in the laboratory chain process. Reasonable quality management requires knowledge about the level of quality that is needed. It is useless to implement and adhere to too strict control rules because this may cause unnecessary, false rejection of assay runs. Ideally, an adequate control procedure should be based on a definition of quality requirements weighing acceptable error against needed clinical decision levels.

5.1. Internal and External QC

Every biomarker assay should include control sample procedures to check the validity of the unknown sample results. Control samples and comparison of their results against control limits should always be integral part of a complete assay procedure.

5.1.1. QC SAMPLES

QC samples are stabilized specimens and available in liquid or lyophilized form because freshly collected sample materials are not always available and unstable for long-term QC use. Important requirements of QC preparations are that they should be time and temperature resistant with little or no vial-to-vial variation, homogeneous, similar in matrix structure to the test material, available at concentrations that cover the physiological range expected in the experimental material, and available in sufficient quantities. Unfortunately these requirements are not always achieved. For serum assays, large pools of serum can be established, aliquoted, and made available to laboratories. However, many manufacturers nowadays supply reference samples on a non-serum–based matrix, and in some cases this yields assay results different from those of true native serum samples. Thus, control samples should resemble as close as possible the analyte fractions representative of those routinely encountered in patient specimens.

5.1.2. MONITORING OF DAILY PERFORMANCE

At least two samples of different concentrations of control material should be included in each assay run to make multirule/decision control procedures possible, for example, by applying Westgard evaluation rules for internal OC (IOC) (9). Thus, repeated measurement of control samples allows to determine imprecision of the assay system. In addition to the use of IOC for day-to-day assay monitoring, the long-term trend in assay performance should be regularly checked in order to detect any shift or drift. Obviously, there should be agreed criteria for batch rejection. Levey-Jennings charts (10) are practical tools to evaluate the controls simply by plotting the individual values on a chart and compare these with a predefined mean with signaling limits (e.g., ± 2 SD). The chart patterns bring different kinds of technical problems (random error, systematic error, etc.) to light, and are also useful, simple tools for investigators or supervisors to decide whether or not assay results are within (or beyond) acceptable ranges and whether the data can be reported. Lot-to-lot variation errors of commercial reagents can be reduced by prescreening of critical reagents and be rejected before use if not consistent.

For external QC (EQC) purposes, preparations distributed by a reference laboratory should be included in assay runs if available. In proper EQC programs, the obtained data of control samples should be submitted to an external organization for statistical evaluation. These programs serve to monitor long-term assay performance within each participating laboratory. Moreover, they provide comparison of assay results between laboratories and between different assay designs or brands, if available. This enables the organization to assess systematic errors between laboratories just by comparing the reported mean values of the individual laboratories with the mean of the total or reference group (all laboratory trimmed mean).

5.1.3. EXTERNAL QUALITY ASSESSMENT (EQA) DOES NOT COVER ALL PROCESS STEPS

EQA based on lyophilized tumor tissue extracts or blood specimens does not allow any conclusion with regard to preanalytical, methodological issues such as variation in tissue collection, transport from operating theater to the laboratory, sample storage conditions, homogenization of tissue, and extraction procedures, as the use of external controls covers only reproducibility of the analytical assay procedure and subsequent computation of data. Providing proper instructions and careful observation of the results obtained is the only feasible way to monitor (between-hospital) variations in sample treatment conditions. Because most clinical trials are carried out on a multicenter basis, the interlaboratory QC is very important but the obtained deviations are most probably underestimations of true differences. Therefore, all steps in the procedure from taking biopsies to reporting assay results to the clinician including the preanalytical items should be conducted according to strict protocol guidelines.

5.2. Normalization of Assay Results

Long-term QA trials on steroid hormone receptors (estrogen receptors [ERs] and progestin receptors [PgRs]) assays by the Receptor and Biomarker Group of the European Organisation for Research and Treatment of Cancer (RBG EORTC) have shown that even highly experienced laboratories, with excellent intralaboratory between-run performance, can have difficulties in directly comparing their results with those of another institution. As variation among laboratories in general appeared to be not random (11), a high interlaboratory coefficient of variation (CV) does not necessarily mean inconsistencies in performance of all individual laboratories. These systematic differences in ER and PgR test results pave the way for calibration (*see* Fig. 1). However, normalization can be achieved only when a marker is homogeneous with only one molecular form present. The presence of more molecular forms of the analyte will yield a broad range of data, especially when different immuno(metric) assays (with varying sets of antibodies, each with other affinities to these molecular forms of the analytes) are used.

6. CONCLUSIONS

An important issue in applying (pre-)clinical immuno(metric) testing kits is that different kits in many cases generate different assay results in the same tumor specimen owing to variation in test design, antibody specificity

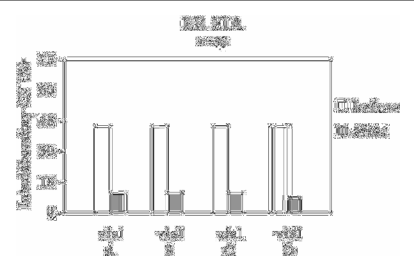


Fig. 1. Interlaboratory CV before (*open bars*) and after normalization of the observed values using a common fifth QA vial (Vial E, gray bars) in estrogen receptor enzyme immunoassay (EIA). Normalization substantially reduces the interlaboratory CVs from 45% to <15%.

and affinity between different kits (or even within a kit from one manufacturer between lots or batches), and use of different calibrators. Although of potential interest, newly explored biomarkers in our view should therefore not be included in large clinical studies unless the assay procedures are carefully evaluated, and common assay protocols, common standards, and QC preparations allowing proper EQA established. At first, such a parameter should be examined in a single expert laboratory. In addition, we strongly advice that in multicenter studies the laboratory performance to be scrutinized prior to generating results from patients in clinical trials.

Hayes proposed criteria for implementing biomarkers in clinical practice, and he defined levels of evidence (LOE) and levels of utility (1). For the highest level (LOE-1) large consistent meta-analysis and validation in a prospective clinical trial should be conducted. Recently, in case of urokinase-type plasminogen activator (uPA) and its inhibitor PAI-1 the level of evidence type-1 was reached, based on the results of a prospective randomized node-negative breast cancer therapy trial (12) and a metaanalysis combining most of the published data sets (13). The therapy trial was under strict external QC by the Receptor and Biomarker Group of the EORTC. The participating laboratories received meticulous instructions on how to run the assays, participated in workshops, used common assays, and were subject of EQC (14). They thus can be regarded as experienced and qualified, which most likely contributed to the success of this trial as well.

Although considerable progress has been made for some analytes as exemplified above for uPA and PAI-1, standardization of biomarker assay protocols, and development of proficiency testing programs for biomarkers, should be an ongoing process. Only the stringent application of QC systems enables a consistent assessment of the prognostic and/or predictive power of biomarkers.

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Tissue Microarrays

State of the Art, Pitfalls, and Promises

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CONTENTS

INTRODUCTION EVOLUTION OF THE TISSUE ARRAY METHOD WHICH ARRAY METHOD AND IN WHAT DENSITY? HOW MANY CORES PER CASE ARE ADEQUATE? CONSTRUCTION OF THE ARRAY IN THE MULTICENTER TRIAL SETTING ANALYSIS SYSTEM SOME UNTAPPED APPLICATIONS OF TMA LIMITATIONS AND FUTURE PERSPECTIVES APPENDIX: AVAILABLE TISSUE ARRAYS REFERENCES

SUMMARY

In this chapter various methods for tissue array construction are compared and protocols for constructing tissue arrays in multicenter trial setting are provided. Listing of all available arrays for breast cancer including commercial and public sources are provided in the appendix.

Key Words: Cancer; multicenter clinical trial; tissue array; tissue microarray.

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1. INTRODUCTION

Although it is possible to examine multiple markers on individual cases by serial sectioning of paraffin blocks, at some point it becomes too costly and time consuming. Tissue microarray (TMA) solves this problem of throughput. Once constructed, assay cost is reduced by more than 100-fold, so screening of multiple markers becomes a reality. Screening 100 cases for 70 markers will require staining and reading of 7000 individual slides, whereas tissue array will reduce this to only 70 slides.

Some of the obvious advantages of using TMA include:

- 1. Large number of cases assessed simultaneously for numerous markers.
- 2. Reduced amount of archival tissues.
- 3. Reduction in cost and time.
- 4. A large number of TMA sections containing different types of tissues, such as a panel of normal tissues, tumors, xenografts, or cell lines, can be produced for testing and optimization of pretreatment conditions, antibody titers, and detection systems.
- 5. Same control tissues can be placed directly on the actual study slides. This helps in ensuring the specificity and sensitivity of immunohistochemistry (IHC).
- 6. Reproducibility of the staining reaction, as well as the speed and reliability of the interpretation, is improved, as all the tissues are on the same slide.
- Consecutive slides can be stained with hematoxylin and eosin (H&E) for morphology or with other antibodies against the same or other molecular targets. This permits comparison of multiple targets in virtually identical, histologically highly controlled regions of the tissues.

As there are many excellent reviews on technical aspects of TMA (1-4), this chapter focuses on topics related to TMA construction in multicenter trial setting and practical guidelines.

2. EVOLUTION OF THE TISSUE ARRAY METHOD

Like any technology, tissue array development has proceeded through almost two decades of evolution (1). In 1986 Battifora described a novel method of creating a block that contains tissue from many cases (5). The creation of this Multitumor (Sausage) Tissue Block involved the tedious process of removing portions of tissue from paraffin blocks, deparaffinization, rehydration to 50% ethanol, and cutting it into 1-mm thick slices and then further into slender rods to produce cross-sectional areas of about 1 mm^2 . These rods were wrapped with small intestine of small animals such as rabbits, and processed into a single paraffin block. The resulting Sausage

Allred (7)	Kononen (1)
Up to 60	Up to 500
2–6 mm	0.6–2 mm
Good	Need at least two to three cores if using 0.6 mm
Skin biopsy punch at under 2 USD (Miltex, York, PA)	Manual arrayer at over 10,000 USD (Beecher Instruments, Sun Prarie, WI)
Large prominent hole evident	Very little damage to the donor block
Easy	Learning curve
	Up to 60 2–6 mm Good Skin biopsy punch at under 2 USD (Miltex, York, PA) Large prominent hole evident

Table 1 Comparison of Two Widely Used Tissue Array Methods

block could be used to screen tissue specific monoclonal antibodies. However, the poor control over the specific location of each component case inside the Sausage limited the value of this strategy as a tool to screen a cohort with clinical follow-up. In 1990 Battifora et al. improved this approach by creating a checkerboard tissue block method in which tissues are evenly distributed in a checkerboard arrangement, and therefore readily identified by their position in the resulting sections (6). However, this method still suffered from the need to go through the tedious process of deparaffinizing the samples as well as the requirement for custom construction of specimen molds with grid pattern. In the same year, Lampkin and Allred devised a novel method of creating a tissue array without the need to deparaffinize the donor samples by using a skin biopsy punch of 3–6 mm in diameter, resulting in up to 40 samples per array (7). Further improvement of this method has resulted in a method that used 2-mm diameter skin biopsy punch resulting in an array containing 60 cores. In 1998, Kononen et al. improved this method further by reducing the core diameter to 0.6 mm, using a high-throughput manual device (1). Owing to affordability of the instrument and increased density of the array, the tissue array has become widely accepted as a high-throughput screening method for candidate molecular markers.

3. WHICH ARRAY METHOD AND IN WHAT DENSITY?

Table 1 summarizes comparison of the two most popular arraying methods. While the method developed by Kononen et al. can generate much higher density array, in reality it may be unnecessary to put so many cores into one array block. Design of an array with more than 100 cores often results in the need for creating subarrays or subsections within the array, which creates problems when navigating between the cores, especially when using fluorescence markers. However, the Kononen method may be preferable to the Allred method because the latter leaves a large hole in the donor block. This defect may create legal or political problems with the pathology department from which the block was provided.

3.1. Beecher Instruments Manual Arrayer

The original commercialized version of the method described by Kononen et al. has popularized the use to tissue array (1). This simple device uses a micrometer to move the recipient block in a precise manner. It has two vertical arms with 0.6-mm punches. The donor punch is slightly bigger in diameter so that the tissue core squeezes into the smaller sized hole in the recipient block. The cores are seeded slightly above the recipient block surface and then later pressed, after incubation at 37° C, so that the height of the seeding can be adjusted to an even surface level.

3.2. Improvements on Beecher Arrayer

When creating a high-density array, the center of the block tends to bulge significantly owing to the increased volume (the diameter of the donor core is slightly larger than that of the recipient hole). In his excellent review of the tissue array technique, Jensen has suggested that keeping the recipient block at approx 100°F using two heating strips attached to the recipient block holder will prevent the problem (2). This compression effect can be resolved, which greatly improves the quality of the array.

3.3. Advanced Tissue Arrayer (Chemicon International, Temecula, CA)

The Advanced Tissue Arrayer device has one advantage over the Beecher Instrument arrayer—adjustable locking Z-height ensures that the top of every tissue sample is delivered precisely to the top of the array block. However, this instrument costs more than five times the Beecher arrayer, and a well-trained histologist or operator will not have problems with height adjustment using the Beecher arrayer. As with the Beecer instrument, during high-density seeding, the center of the block will bulge and require pressing.

3.4. Automated Arrayer ATA-27 (Beecher Instruments, Sun Prarie, WI)

The Automated Arrayer ATA-27 is a high-throughput system that generates three replica arrays containing one core each from 100 cases in about 4 h. Based on a precision XYZ robot, it allows convenient walkaway mapping on a video screen. The only drawback stems from its small footprint that allows loading of only 20 cases at a time. Theoretically the array generated using this automated system should be much superior to manually constructed ones owing to its Z-axis control—that is, control of the depth of seeding of cores into the array block. However, because it also creates a compression effect, this potential advantage is lost. Although one could heat the recipient block, as described by Jensen for manual arrayer, the block holding mechanism and moving parts in the ATA27 make it difficult to attach a heating strip. One solution has been to heat the entire enclosed environment to 37°C using an air heating system. However, temperatures higher than this cause the donor blocks to soften and the tissue core push through the block.

4. HOW MANY CORES PER CASE ARE ADEQUATE?

A potential drawback of testing a marker in a TMA is that a small core may not be representative of the whole tumor, given the likelihood of heterogeneity. Heterogeneity of the marker expression may not be captured if only one core is sampled especially for 0.6-mm core sampling. Camp et al. examined the correlation between cores and whole sections and came to the conclusion that sampling of two cores are sufficient for breast lesions (8). Zhang et al. found even one 0.6-mm core adequately represents the whole section (9). However it is a safe practice to sample three cores from random spots of each tumor case. In our practice, we routinely sample three cores and seed them into three separate arrays. For initial marker screening we use one core. Promising markers are then validated using all three cores.

5. CONSTRUCTION OF THE ARRAY IN THE MULTICENTER TRIAL SETTING

5.1. Handling of Donor Blocks— Cutting, Staining, and Mapping of the Region of Interest

The standard operating procedure (SOP) for TMA construction at the National Surgical Bowel and Breast Project (NSABP) Division of Pathology Laboratory is as follows:

- 1. Align blocks to microtome knife with Histo-Collimator (Richard-Allan Scientific).
- 2. Cut one tape section with Paraffin Tape-Transfer System (Instrumedics).
- 3. Stain with H&E and map the region of interest/core sampling target with red Sharpie permanent marker.
- 4. Overlay the H&E slide to the original block to mark the core sampling target with red Sharpie permanent marker.

- 5. Sort the blocks according to the thickness of the tissue in the block.
- 6. Recipient array block is heated to 100°F before arraying.
- 7. Construct the array with three spleen cores as position marker on one corner.
- 8. Seed 100 cores in each array block containing one core from each 100 cases. Three replica arrays are generated.
- 9. Incubate the array block at 37°C for 10 min before pressing with the glass slide to level the surface of the array block.

Alignment of the block surface to the microtome knife can be quite varied, posing a serious problem to the histologist. Significant time may be wasted orienting the block to the knife. More importantly, several sections of tissue may be lost before a good-quality section can be cut. Use of a microtome equipped with a Histo-Collimator (product no. 755130. Richard-Allan Scientific, Kalamazoo, MI) can largely eliminate this problem. Using an optical alignment method, the Histo-Collimator allows correct alignment of the block surface to the angle of the microtome knife.

Mapping of the region of interest, especially small areas such as normal lobule, can be difficult and often inaccurate when using routinely processed H&E stained sections, because fatty breast tissue expands or becomes deformed in the water bath before the section is picked up on the glass slide. To solve this problem, we have been using the Paraffin Tape-Transfer System (Instrumedics, Hackensack, NJ) that bypass the water bath step completely. First, a plastic tape is attached to the block surface and a section is cut. The section is mounted on the tape and laminated on a special slide coated with UV curable acrylic, which is cured by brief exposure to UV light. The tape is then removed using an organic solvent and the slide is stained with H&E. This procedure will produce an H&E slide that is an exact match to the original tissue in the paraffin block allowing precise mapping of the region of interest for core sampling. We use red colored permanent marker pen to mark the region of interest for two reasons. First, it is readily identifiable for sampling and second, after core transfer to the array block and during sectioning, the red colors on the core provide visual landmarks to evaluate the quality of the array. By combining Histo-Collimator with Paraffin Tape-Transfer System, the mapping of the donor blocks can become a highly efficient process.

5.2. Handling of Refusal to Submit Blocks

The major benefit of TMA construction is to facilitate analysis of large numbers of tissues while preserving time, money, and tissue resources. Thus, TMA preparation is ideal for tumor marker studies in large cooperative groups, such as the NASBP. However, in such an organization, individual institutions may be reluctant to share paraffin blocks. One solution is to provide an alternative to block submission. For example, NASBP accepts submission of a 2-mm core plus a minimum of 10 unstained sections as an alternative to block submission. This strategy has received favorable feedback from the membership. A skin punch biopsy device used for the Allred method (Miltex 2-mm skin biopsy punch) is provided to each member institution on request and the device is shipped back after sampling the tissue without the core extraction from the device to minimize the end user time and effort. We create Allred arrays from these cores, cut and H&E stain, map it, and resample three 0.6-mm cores from them.

6. ANALYSIS SYSTEM

Several systems have been developed to aid in navigation and scoring of assay results for tissue microarray. Table 2 summarizes some of the commercial systems that can be used for navigation. Depending on the need of the institution or investigators, there are wide variety of system choices. At this point ACIS (ChromaVision, San Juan Capistrano, CA) seems to provide the most tested algorithm for immunohistochemistry (IHC) marker analysis. This system has the advantage of being available in most large pathology departments in the United States. Access and image analysis can be accomplished through Web interface systems, such as the BLISS system (Bacus Laboratories, Lombard, IL) and ScanScope (DakoCytomation, Glostrup, Denmark).

Kemp et al. have described an image analysis system that is based on double-immunofluorescence staining with the marker of interest and epithelial cell markers. In this system, tumor epithelial cells can be automatically detected, and therefore staining intensity of only the epithelial cells is measured quantitatively. This has resulted in scoring that has a wider dynamic range than the usual IHC analysis (10).

7. SOME UNTAPPED APPLICATIONS OF TMA

7.1. Use in the Clinical Laboratory

In a typical reference laboratory setting, batch processing of clinical cases using a tissue array would be quite possible. In this case, the reference laboratory might conduct a new TMA periodically (e.g., once a week), prepare slides for IHC for the markers of interest, analyze them, and report the data back to the individual caregiver for routine clinical use. The remaining TMA would then be available for future research studies. Such a strategy is currently employed in some cancer agencies regional laboratories in Canada for immunohistochemical markers such as estrogen receptor (ER) and for fluorescence *in situ* hybridization (FISH) for HER2. While objectivity and reproducibility of the IHC could be questionable, studies have

	Con	nmercially Available	Commercially Available TMA Analysis Systems	JS		
System	Hardware	Mapping of cores	Image analysis	Fluorescence capability	Remote viewing through Web interface	Remote image analysis through Web interface
BLISS (Bacus Laboratories, Lombardi, IL)	Remote pathology station based on motorized microscope and Web slide server	Semiautomatic, interactive Each core image is an montage of six image files	Has proven track record of developing CAS 200 system	No	Yes	Yes
ACIS (ChromaVision, San Juan Capistrano, CA)	Motorized microscope based image analysis station Auto slide feeder	Semiautomatic, interactive	Has proven track record for HER2 IHC and CISH scoring	No	No	No
ScanScope (DakoCytomation, Glostrup, Denmark)	Remote pathology station, scanner based	Semiautomatic, U interactive Can save individual core images as TIFF or JPEG file format for exporting	Under development L Lat	No	Yes	Under development
ARIOL (Applied Imaging, New Castle Upon Tyne, UK)	Motorized microscope based image analysis station Auto slide feeder	count programs Semiautomatic, interactive	High-throughput image analysis system	Yes	No	No
Icys Research Imaging Cytometer (Compucyte, Cambridge, MA)	Laser scanning cytometer	Mapping using low power scan mosaic	Dedicated three- channel laser scanning of three fluorescent dyes Highly quantitative	Yes	No	No

Table 2 Commercially Available TMA Analysis Systems

demonstrated the feasibility of improving this with pixel-based image analysis system such as ACIS from Chromavision.

7.2. Survey of Amplicons

One interesting potential use of tissue microarray is in combination with array-based comparative genomic hybridization. Recently Pollack et al. (11) and Hyman et al. (12) have conducted a comprehensive survey of amplifications and deletions in the breast cancer genome and correlated the results with gene expression levels using the same cDNA microarray chips. Both groups found significant contribution of gene amplification in the transcriptional activity of the cancer genome. These genes are obviously good candidates for prognosticators as well as therapeutic candidates. Hyman et al. have examined one of these amplified genes *HOXB7*, by FISH on tissue microarray and found it to be prognostic (12). However, they did not go further to screen all of the described amplified and overexpressed genes and screen them rapidly on tissue microarrays. FISH is in general more objective than immunohistochemistry because it relies on spot counting—therefore such markers may turn out to be clinically useful ones.

8. LIMITATIONS AND FUTURE PERSPECTIVES

When the tissue array method was first introduced, it was expected to revolutionize the development of cancer marker studies. However, progress has been slow, for at least two reasons: (1) difficulty and high cost of creating antibodies that work for immunohistochemistry and (2) lack of availability of tissue resources with solid clinical outcome data. One effort to resolve the latter problem is the Cooperative Breast Cancer Tissue Resource (CBCTR), established by the National Cancer Institute of the United States. Based on distributed tissue banks of paraffin blocks in the United States, CBCTR not only provides individual tissue sections but also makes tissue microarray arrays of various designs from breast cancer available. One of the most valuable arrays from CBCTR has been the "progression array" containing cores from various lesions through the presumed stages of breast cancer progression. However, the CBCTR has not developed tissue arrays for prognostic or predictive marker studies. Recently the NSABP has decided to release tissue microarrays generated from NSABP clinical trials to the scientific community, using NCI as an honest broker. Availability of the first array containing 2000 cases from NSABP trial B-28, in which nodepositive breast cancer patients who all received adjuvant doxorubicin and cyclophosphamide (AC) were randomly assigned to receive paclitaxel or not, was announced in 2003. While individual laboratories may not be able to develop IHC for 100 different markers at once, 100 laboratories in the world perhaps can share this burden and achieve the same aim. It is hoped that other clinical trial groups and banks will follow similar steps to make their resources as widely as possible.

9. APPENDIX: AVAILABLE TISSUE ARRAYS

9.1. Commercial Sources

As listed in Table 3, many companies now offer both multiorgan site arrays as well as breast cancer specific arrays.

9.2. CBCTR Tissue Array (http://www-cbctr.ims.nci.nih.gov/)

This is currently the best resource to investigate the role of a candidate marker on breast cancer progression once the assays have been optimized for paraffin sections and arrays using in-house or commercially available arrays. The National Cancer Institute (NCI) Cooperative Breast Cancer Tissue Resource (CBCTR) is funded by the NCI to supply researchers with primary breast cancer tissues with associated clinical data. The CBCTR has designed a breast cancer TMA that can be used to investigate differences in prevalence of potential markers in three stages of invasive breast cancer: node-negative, node-positive, and metastatic disease. All of the invasive cases are primary breast cancers with a principal histology of ductal cancer accessioned through the CBCTR. The arrays were designed by National Cancer Institute statisticians to provide high statistical power for studies of stage specific markers of breast cancer.

Each TMA block consists of 288 0.6-mm cores taken from paraffinembedded specimens that represent 252 breast cancers and normal breast specimens plus 36 controls. Each array is created in quadruplicate to address possible tissue heterogeneity: four cores are taken from each specimen block, with one core per specimen appearing in each of the four replicate array blocks. The information provided for each case on the array are: tumor size, Tumor–Node–Metastasis (TNM) stage, number of nodes positive, grade, age at diagnosis, and race.

The 252 normal breast and breast cancer cores appearing on each TMA block include:

NODE-NEGATIVE BREAST CANCER: 64 cores.

NODE-POSITIVE BREAST CANCER: 64 cores.

METASTATIC BREAST CANCER: 64 cores.

DUCTAL CARCINOMA IN SITU (DCIS): 20 cores (10 from individuals without an invasive disease component and 10 from individuals with invasive disease represented elsewhere on the TMA).

NORMAL BREAST TISSUE: 40 cores (20 from individuals without breast cancer and 20 from individuals with breast cancer represented elsewhere on the TMA).

		List of 1 MAs t	List of TMAs from Commercial Sources		
Company	Website	Product	Core	Clinical data	Composition
Ambion	http://www.ambion.com	LandMark high-density 260 Cases breast tissue microarrays	260 Cases	Stage, surgery type, tumor size, node status, histologic type, age, and sex	Mixed infiltrating ductal carcinoma, lobular carcinoma, and matched benign tissue
Clinomics	Clinomics http://www.clinomicslabs.net	LandMark low-density 72 Cases breast tissue microarrays HD-BR-1 200 Case	72 Cases 200 Cases	0	
	-	L.D-BR-1	60 Cases	Complete pathology data, medication history, outcome, and follow-up data	Representing virtually all stages of breast cancer, from DCIS through stage IV, includes matched normals, benign disease (e.g., fibrocystic disease), and clinically defined
Biogenex	www.biogenex.com	High-density tissue microarray	200 (0.6-mm diameter) Age, sex, organ, morphology, tumor size, ori (biopsv/surger)	Age, sex, organ, morphology, tumor size, origin (biopsv/surgerv)	normats.
		Low-density tissue microarrav	50 (1.2-mm diameter)		
Chemicon	Chemicon http://www.chemicon.com	Chemicon select tissue array	6 (2-mm diameter)	N/A	Four cancer, two normal tissue
					(continued)

Table 3 List of TMAs from Commercial Sources

	Composition	Tumors are different histological grades from various stage groups; 3 Stage I, 13 stage II, 14 strate II, 10 strate IV	Tissue cores 1-2 and 4-49 are infiltrating ductal breast adenocarcinoma, tissue core 3 is breast mucinous carcinoma, tissue cores 50-54 are medullary carcinoma, tissue cores 55-57 are ductal carcinoma in situ (DCIS), tissue cores 58-59 are tubular breast cancer, and tissue core 60 is infiltrating nodular odenocreitoronodular	Mixed breast cancer, fibroadenoma,	Σų
Table 3 (continued)	Clinical data	Associated clinical data	N/A	N/A	N/A Age, sex, organ, diagnosis, T stage, LN, ER, PR, p53
	Core	50 (1-mm diameter)	60 (1.5-mm diameter)	70	60 (2mm) 60
	Product	Standard breast cancer array	MaxArrayTM breast carcinoma tissue microarray	Receptor Grid TM tissue array—breast	CB1 IMH-304 (CB) human, breast cancer
	Website	http://www.ardais.com	http://www.zymed.com	http://www.abcam.com/	http://www.gentaur.com/ http://www.imgenex.com/
	Company	Ardais	Zymed	Abcam	Gentaur Imagenex

The 36 control cores appearing on each TMA block include:

NORMAL NON-BREAST TISSUE: 16 cores (4 cores from each of the following tissue types: kidney, endometrium, prostate, and appendix). CELL LINES: 20 cores (5 cores from each of the following cell lines: HT-29, PC-3, MCF-7, and T-47D). These cell lines have previously been characterized with regard to expression of a variety of markers of interest in breast cancer.

9.3. NSABP

(National Surgical Adjuvant Breast and Bowel Project): (http://www.nsabp.pitt.edu)

NSABP conducts large phase III clinical trials and started offering tissue arrays constructed from its trials to general scientific community through NCI. The first available array is constructed from 2000 cases enrolled in its trial B-28, which addressed the question of adding sequential paclitaxel to four cycles of adriamycin plus cyclophosphamide. NSABP plans to offer tissue arrays from other protocols as they are constructed. These arrays are great resources for studying marker-by-treatment interaction questions, although they suffer from being inherently underpowered.

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II Assays for Gene Expression and Post-Translational Protein Abnormalities

4

Gene Expression Profiling with DNA Microarrays

Revolutionary Tools to Help Diagnosis, Prognosis, Treatment Guidance, and Drug Discovery

Michele De Bortoli, PhD and Nicoletta Biglia, MD

CONTENTS

INTRODUCTION How a Microarray Experiment Works: The Technical Principle DNA Microarray Analysis of Breast Tumors Current Debates References

SUMMARY

DNA microarrays are small solid supports on the surface of which DNA probes for thousands of genes have been orderly arrayed. Hybridization of labeled RNA from tissues or tumors allows evaluation of the relative amount of any specific mRNA present in the samples, depicting its gene expression profile. During development and progression of breast cancers, specific genetic programs are activated, which can be assayed on DNA microarrays. Studies performed

Cancer Drug Discovery and Development: Biomarkers in Breast Cancer: Molecular Diagnostics for Predicting and Monitoring Therapeutic Effect Edited by: G. Gasparini and D. F. Hayes © Humana Press Inc., Totowa, NJ so far show that the gene expression profile of a tumor defines its biology, its invasive and metastatic potential, and its responsiveness to treatments. Classification of breast cancer by expression profiling appears a very powerful approach, outperforming all commonly used methods of classification. However, much work has to be done before microarray will give robust information for clinical decisions and become routine practice.

Key Words: DNA microarrays; gene expression profiling; breast cancer.

1. INTRODUCTION

A modern view of genetics is that the genomes of all organisms, most spectacularly those of Metazoa, are made of—rather than genes—genetic programs, that is, ensembles of genes that coordinately specify developmental phases, cell types, tissue and organ morphogenesis, and cellular responses to endogenous and exogenous stimuli.

The genomic era has led to the complete description of the DNA sequence of a number of organisms, from *Escherichia coli* to humans—in other words the complete chemical structure of genetic materials, represented as huge strings of A, T, C, and G, stored and accessible in public databases (*www.ncbi.nlm.nih.gov*, to cite the most popular). Saying that we know the complete DNA sequence of some organisms does not mean that we know what the sequence means, even though, at an increasing rate, DNA sequences in databases become annotated with functional information. Genome sequencing was one fundamental step to move on to functional genomics, that is, to understand the functions of all the genes, to unreveal regulatory networks and perhaps to discover many unsuspected genetic languages and functions.

DNA microarrays are tools to analyze genetic programs. Hybridization of the entire messenger RNA (mRNA) pool expressed by a tissue to DNA microarrays, containing probes to theoretically all the genes of an organism, allows to evaluate in a single analytical step which genes and to which extent they are transcribed, that is, in technical terms, the gene expression profile. This gives an immediate and complete picture of genome activity in a specific biological situation, or an estimate of the difference between two biological situations. For example, one of the first published applications of DNA microarrays was the analysis of genome activity changes during metamorphosis in *Drosophila (1)* or during sporulation in yeast (2). These kinds of experiments give remarkable results and open the way to completely new studies, for example, to analyze regulatory sequences of coregulated genes to discover common regulatory pathways, to find the coordinated actions by which cells respond to pathogens or drugs, or to describe changes in genomic

structure and expression that accompany tumorigenesis and cancer progression. Several articles, indeed, have reported gene expression profiles of human and experimental tumors and provided exceptionally interesting, albeit preliminary, evidence on the genetic reprogramming during cancer development, progression, and response to treatments.

This chapter deals with the use of gene expression profiling in breast cancer, showing that it will represent an unprecedented powerful tool for reclassifying breast cancers to formulate prognosis, predict and monitor response to treatments, and provide information on the pathogenetic mechanisms relevant to designing new therapeutic interventions. At the same time, we will try to make the focus on the number of steps that are necessary before applying these tools to patient care.

2. HOW A MICROARRAY EXPERIMENT WORKS: THE TECHNICAL PRINCIPLE

Genomics and microarrays have pervaded the scientific literature so extensively during the past 5 yr that probably no further description is necessary. Some excellent reviews have been published (3-6). In brief, a microarray is a physical support (nylon filters, plastics, glass slides) presenting on the surface an ordered array of hundreds or thousands of DNA sequences, representing probes (i.e., the complementary sequence) for each mRNA species, that is, primary gene products. To produce a gene expression profile, the RNA is extracted from tissues or cells, labeled, and hybridized to the array. Hybridization, which is highly specific, produces labeled spots whose intensity is proportional to the level of expression of that particular sequence (gene) in the sample. In the very first applications of this methods, nylon membranes were spotted with one to a few hundred cDNA probes using vacuum manifolds or spotters and hybridization of radioactively labeled RNA was evaluated by densitometry. Evolution of this technique has produced DNA microarrays or DNA chips, usually made of microscope glasses on which cDNA or oligonucleotide probes representing up to 50,000 human genes are orderly arrayed. Fluorochrome-labeled sample RNA hybridizing to the array gives fluorescent spots that are quantitatively measured by laser scanning.

2.1. DNA Chip Technology

Today there are three main alternative techniques for microarray production and two principal approaches for hybridization. The historically first approach is represented by the robotic spotting of probes onto the surface of microscopic glasses. This was introduced by Pat Brown's laboratory (7) and is widely used at present both by biotech companies and in university facilities. Probes are usually cDNA fragments (200–1000 bp) produced by polymerase chain reaction (PCR) from collections of clones or—more recently—chemically synthesized oligonucleotides, 50–70 bases in length, representing relevant parts of the genes. These chips may accommodate up to 50,000 spots/cm². The second approach, which was pioneered by Affymetrix, makes use of oligonucleotide probes, 20–25 nt in length, that are directly synthesized on a microscope glass using a photolithographic technique, at a very high density (up to 300,000 different oligos/cm²) (8). Third, chemical synthesis of oligonucleotides, up to 70 nt in length, on solid supports is realized, at densities of up to 50,000 different sequences per slide, with an ink-jet technology from Rosetta Inpharmatics (9).

2.2. Samples

Sample preparation, always based on the production of copies of mRNA (cDNA) by reverse transcriptase enzymes, also presents alternatives. The most popular method is the dual-color fluorogenic technique, in which two different fluorochromes (most often the cyanines Cy3-red and Cy5-green) are incorporated in cDNA during or after reverse transcription, the first in the interrogated sample and the second in a reference sample (e.g., RNA from tumor labeled with Cv3 and RNA from normal tissue labeled with Cy5). The two samples are then cohybridized to the microarray and a twocolor image is rescued, in which the ratio between colors directly reflects the ratio of abundance of any particular mRNA species in the sample vs control experiment, that is, it gives a measure of differential expression. In alternative, a single-labeled RNA is hybridized to the array. This is the case of radioactive labeling, when using DNA arrays made on nylon or plastic membranes, or when using short oligonucleotide probes, as in the case of Affymetrix chips. In this case, incorporation of a small label (e.g., biotin) during RNA copy production is preferred, followed by posthybridization fluorogenic staining, for example, with fluorochrome-conjugated avidin.

Significant technical advancements were achieved toward nanoscale sample preparation. In fact, the main limitation for microarray analysis is the requirement of relatively large amounts of good-quality RNA. Techniques of linear amplification were developed that allow sample preparation, good for microarray hybridization, from as little as 10 ng of RNA (10). A T7 promoter sequence is attached at the 5'-end of the first cDNA strand during reverse transcription, then the final double-stranded cDNA template is used for in vitro transcription by T7-RNA polymerase, producing several copies of complementary RNA (cRNA). This method was shown to amplify the mRNA in a quasilinear fashion, thus allowing gene expression profile analysis in very small amount of tissue, as, for example fine-needle biopsies or laser-dissected tissue specimens (11,12).

2.3. Data Analysis

The primary outcome of a microarray experiment is a fluorescence image. Spot quantitation produces a long list of hybridization values that needs interpretation. The computational analysis of these results is a very important part of the story, and in parallel with technical advances in microarray production and hybridization methods, several computational tools and packages have been developed, allowing spot quantitation, background subtraction and normalization, data storage and retrieval, and higher-level analysis, that is, statistical as well as clustering analysis. Statistics is applied to measure the significance of variations seen among different samples analyzed (e.g., cells treated with the drug X in triplicate vs triplicate untreated controls). In the case of microarrays, considering each gene as a single entity and applying common statistical difference tests would mean losing much of the information that combinatorial analysis can provide. Thus, dedicated statistics should be developed, one example of which is the widely used and publicly available SAM analysis, and further developments (13). The second step is represented by clustering analysis, that is, the search for genes that are coregulated in a given biological situation and/or samples that show similar gene expression patterns. There are many different methods to produce clustering, but in any case a measure of similarity is used to place each gene (and/or experimental point) into a single group or a rank within a hierarchy. Outputs are ordered lists of value but are more commonly displayed as a two-color matrix (14) using false-color scales to show different levels of expression or sample/reference expression ratio.

One important aspect of data analysis is data validation which, at least in the studies conducted so far, has been systematically performed by analyzing either the mRNA levels of selected genes among those resulting from microarray analysis, using established methods, such as real-time quantitative reverse transcriptase-PCR (RT-PCR), Northern blotting or RNase protection assay, or by immunohistochemistry/*in situ* hybridization. It is very important to note that in all published studied, a general consistency of expression data obtained with microarrays was found, demonstrating the general robustness of this technique.

3. DNA MICROARRAY ANALYSIS OF BREAST TUMORS

Gene expression profiling in breast cancer is being used to address a number of questions, such as which genes are activated (or down-regulated) during progression through the different stages, or which are the genes that are associated with metastatic potential, which are those that mark responsiveness to certain treatments (such as antiestrogens), and so forth. One preliminary and important question was: Because breast cancer is characterized by a high level of heterogeneity, can gene expression profiles help classify cases in a clinically useful fashion? Preliminary studies showed, in fact, that different breast carcinoma-derived cell lines display distinctive gene expression patterns under in vitro manipulation and that those patterns can be recognized in human breast tumors (15).

The first study, performed with a sufficiently wide microarray (8102 human genes) and with a reasonably large number of breast tumors, confirmed this hypothesis and gave many important insights (16). This article by Perou et al. was entitled "Molecular portraits of human breast tumours" to suggest immediately that a gene expression profile is a multicomponent characteristic that is *individual* but at the same time allows recognition of similarities among samples, exactly as we do when we recognize people and discriminate members of the same family by resemblance. This is an essential property of microarray analysis, that is, results are valid per se, independently of the function of every single gene that compose the profile, in the same way as recognizing people in most cases does not require concentrating on any single character, but just appreciating (integrating) the sum of all the characters. The breast tumor specimens analyzed in this study, in fact, gave individual profiles of expression, but easily recognizable as belonging to well-defined groups. First, in the study there were 22 coupled samples, that is, taken from the same patients, either primary tumor vs lymph node metastasis or tumors biopsied before and after doxorubicin chemotherapy. This allowed a remarkable conclusion: the tumors from the same individual were always much more similar to each other than to any sample from different patients. Second, tumors could be grouped together on the basis of similarity of their expression profiles to specific cell lineages, that is, luminal, basal, and myoepithelial cells. These observations, confirmed in successive studies, suggest that cell lineage, as related to the clonal origins of a cancer, is the fundamental determinant of the expression profile. Individual tumors are the result of microevolutionary processes, during which mutations in the genome have been selected by the environment ("host"), rendering the cells able to escape "normality," that is, the number of controls and checkpoints that normal cells in a normal organism exert toward each other. As in all evolutionary processes, there is no particular order in the history of genetic alterations and selection steps, so that similar end points can be attained following different pathways. This is reflected, in clinical terms, by the differences shown by individual cancers, in terms of histological appearance, local advancement, metastatic capability, time to relapse, sensitivity to chemical treatments, and so forth. In other words, it is conceivable that each tumor constitutes an individual entity from genetic and molecular aspects, exactly as members of a population do, and this underlies clinical variability.

3.1. Pathogenesis and Progression

As mentioned earlier, the epigenetic point from which tumor cells start to deviate (i.e., the cell lineage) remains recognizable as such in the expression profile. This has been clearly demonstrated also for other types of cancer, such as leukemias and liver cancers (see ref. 17 and references therein). In other words, gene expression profiles can "trace" the evolutionary history of a tumor. This point has been reinforced by more recent studies addressing the important point whether genome activity changes during progression from atypical ductal hyperplasia to invasive tumors. The group leaded by Dennis Sgroi developed a laser-capture microdissection technique, coupled to RNA amplification, that allowed gene expression profiling using 12,000-gene (12K) microarrays on different sections from the same tumor, showing pathologically discrete stages, in comparison to normal adjacent tissue samples (12). Results showed that the different stages of progression are very similar to each other in the same patient. Furthermore, the different synchronous stages of tumor within an individual patient clustered more closely to one another than to their respective stages from other patients. These data confirm that alterations in gene expression pattern, as compared to normal tissue, are present in the earliest stage of tumor development and are remarkably stable during progression. This implies that the structural changes leading to tumor development imprint in a quite stable manner genome activity in tumor cells, in other words that the genetic point of origin can determine the final expression pattern. This question was directly addressed on the experimental model systems by Desai et al. (18), by examining mammary carcinomas from six transgenic animal lines carrying different oncogenic constructs. This study demonstrated that the gene expression profiles are determined by the signal transduction pathway activated by any particular oncogene. ras, neu, and Polyoma Middle T clustered together since they all converge on the ras/MAPK pathway, while myc or SV40 Large T-induced tumors showed distinct expression patterns. The same conclusion was reached in a study by Shan et al. (19) showing that mammary tumors induced in rat by two different carcinogens, dimethylbenz[a]anthracene and 2-amino-1-methyl-6-phenylimidazol[4,5-b]pyridine, display very distinctive gene expression profiles, notably associated with indistinguishable histology.

Many studies during the last two decades have reported the diverse genetic alterations that are detectable in tumor cells, comprising pointmutations, translocations, loss of heterozygosity (LOH), and amplifications. Some of these alterations are very relevant to breast cancer, for example, amplification of the *erb*-B2 oncogene. Any individual tumor at diagnosis presents a number of alterations, in patterns that until now, and with few exceptions (*erb*-B2, for instance), were not traced to any clinically relevant condition. One important question is whether the pattern of genetic alterations has a major impact on gene expression. cDNA microarray analysis allows this question to be addressed directly. In fact, the DNA extracted from tumors can be fragmented, labeled with one fluorescent dye, cohybridized together with normal DNA from the same donor, and labeled with a second fluorochrome, to the same cDNA microarrays used for parallel gene expression profiling on RNA from the same tumor. Using this approach, Pollack et al. (20) and Hyman et al. (21) were able to show that gene copy number indeed has a major impact on gene expression, at least for a number of loci. The pattern of gene amplification or deletion was consistent with previous cytogenetic studies of breast cancer. In the first study, which was done on 6.7K mapped human genes, it was observed that 62% of the genes showing high-level amplification displayed moderate or elevated expression, and that the range of amplification is generally reflected in the level of expression. Overall, it was found that 12% of all the variation in gene expression can be attributed directly to underlying variation in gene copy number (20). In the second study, using microarrays featuring probes for 14K mapped human genes, it was confirmed that gene amplification or loss had substantial effects on gene expression, identifying 270 genes whose expression level was directly linked to gene copy number (21).

These articles demonstrate that microarray analysis can give very important information on the pathogenetic mechanism and also define genetic alterations in subsets of tumors that, because they produce stable expression changes, represent important potential targets for drug development.

3.2. Classification, Prognosis, and Prediction

In the aforementioned articles, it was quite clear that human breast tumors show distinctly different gene expression profiles that, at the same time, bear resemblance with the cell lineage from which they most likely originated. This does not sound as a completely new concept, as it represents the principle of the common pathological classification. Under the light microscope, indeed, the appearance of a tumor often resembles the normal cell of origin, especially when some immunohistochemical marker is used in combination, such as cytokeratins.

Hierarchical clustering of gene expression profiles leads to a classification of breast tumors that follows in part that resulting from classical pathological or biochemical markers but, importantly, also suggest new categories and new subclassifications. Indeed, it is possible to identify groups of genes whose expression profile, or "signature" correlate with any "dominant" condition of the tumor. For example, the main discriminant in the biology of breast cancer, that is, the presence of steroid receptors, can be easily and clearly distinguished by a "signature" that possibly reflects many estrogenresponsive genes (16,22-24). The same was true also for tumors containing an amplified *erb*-B2 oncogene, which were easily recognized by gene signature (16,20) and linked to several other coamplified genes (25) and to fatty acid metabolism (26) by data mining of microarray results. Specific gene expression signatures were also observed for hereditary breast cancer, with distinct signatures for *BRCA1* and *BRCA2* tumors (27). In addition, microarray profiling was shown very effective in discriminating different groups in non-*BRCA* hereditary cancers, paving the way to discovery of the genetic lesions behind these (28).

The sum of the observations reported in the preceding also justifies the hypothesis that tumors at diagnosis have distinguishable properties characterizing their future behavior. This represents a very important issue, which was addressed in the past, in breast as well as in other cancers, by the use of single biochemical or genetic markers, or based on the clinicopathological data. The fact that tumors with different expression patterns may represent clinically different classes with distinct risks of metastasis can be addressed with retrospective studies. In a further analysis of data already published from a 8.1K microarray analysis of locally advanced breast cancer (16), Perou et al. found that the classes of tumors, previously defined on the basis of gene expression signatures, that is, normal-like, basal-like, luminal-like, erb-B2⁺, could be subdivided into further clusters by limiting the analysis to the most significant 456 genes and applying a new significance analysis procedure (29). Interestingly, these new subgroups showed different frequencies of mutation to TP53 gene and different survival (29). The relatively small cohort of patients analyzed in this study and the fact that most of them were at stage III undergoing different treatments made it difficult to generalize the findings. However, in a more recent study performed using a much larger microarray platform, these finding were substantially confirmed. Van't Veer et al. (22) used microarrays developed at Rosetta Inpharmatics, containing oligonucleotide probes for 25K human genes, to profile gene expression in breast tumors from 117 young patients. A supervised hierarchical clustering algorithm was able to identify a gene expression signature strongly predictive of a short interval to distant metastasis ("poor prognosis" signature) in 78 patients with lymph node-negative disease. Further analysis showed that 70 genes were sufficient to correctly predict the clinical outcome in these patients. On the basis of these results, this analysis was successively extended to a cohort of 295 consecutive patients, comprising both lymph node-positive and -negative cases (30). The "poor prognosis" signature was displayed by 180 patients, with an average overall 10-yr survival rate of 54.6% vs 94.5% (50.6% vs 85.2% for disease-free survival) shown by patients with the complementary "good prognosis" signature. Importantly, these highly significant differences hold true when patients were analyzed according to lymph node status. The prognostic power of gene expression signatures was stronger than any other clinicopathological parameters in predicting disease outcome. This study was the first to demonstrate that breast cancer classification based on gene expression profiling can outperform current classification methods and give concrete help in selecting patients for adjuvant therapy (31). A further issue that can be addressed by microarray analysis is prediction of response to therapies. In a very interesting study, Sotiriou et al. (11) determined 7.6K gene expression profiles on fine-needle aspirates (FNAs) taken from patients eligible for neoadjuvant chemotherapy. The analysis was repeated at surgery and hierarchical clustering not only confirmed the individuality and stability of gene expression profiles (successive FNAs from the same patient or the FNA and surgical biopsy from the same patient always clustered together), but also demonstrated that microarray analysis before treatment can predict the response to chemotherapy. In addition, the authors were also able to identify a small group of genes whose expression changes during chemotherapy, suggesting a new way to monitor response.

Of course, one of the main interests in breast cancer management is to define hormonal responsiveness of tumors, since the presence of estrogen receptors (ERs) and progestin receptors (PgRs) identifies tumors that can benefit from antiestrogenic treatment with limited precision (60-70%). This is a more difficult issue to address, because retrospective, uniformly treated, evaluable cohorts of patients are hardly available or not at all. For this reason, most of the studies concentrated either on the ability of gene expression profiles to identify correctly the ER/PgR status in breast tumors, or focused on "in vitro" model systems, in which gene expression changes following estrogenic or antiestrogenic treatments can be measured. In all the studies on breast tumors reported in the preceding, the main clustering pattern clearly distinguished ER⁺ from ER⁻ tumors, providing further proofof-concept. Conversely, various "in vitro" studies have evidenced large sets of genes whose expression is regulated by estrogen or antiestrogens (32-37). These results must be validated by studying the expression of these genes in patients in whom genuine antiestrogenic response can be assessed, such as in the advanced set. Interestingly, the expression pattern of a handle of genes identified in such an "in vitro" experiment (37) is enough to correctly detect the ER status of breast tumors (24). At the same time, these types of studies may elucidate a number of pathways that are activated in hormone-unresponsive tumors, giving insights into new targets for drug development.

4. CURRENT DEBATES

It is evident that the major advantage of gene expression profiling is given by *combinatorial analysis*. Previous work attempting to define biochemical or genetic markers that could help the clinical management of breast cancer was concentrated on single variables. When facing a complex disease such as breast cancer, featuring great clinical variability and molecular heterogeneity, it is expected that single markers may show only very partial association with tumor characteristics. Gene expression profiling, indeed, takes into account thousands of variables at the same time and singles out the individual groups of genes that associate with a relevant feature: none of them, individually, will show significant association with the parameter considered, but the sum of these does. It is noteworthy that the most important (to date) set of genes found for breast cancer, that is, the 70 genes described by Van de Vijver et al. to discriminate the risk of metastasis (30), do not comprise any of the most popular genes previously proposed as prognostic markers. In this study, the authors started with a microarray featuring probes for 25,000 human genes. Of these, approx 5000 were found with significant variations in at least 3 of the initial 117-tumor cohort. Of these, 231 genes were found associated with clinical outcome and, finally, an iterative exclusion procedure narrowed the number of predictor genes down to 70(30). This is a quite common feature of this kind of study. The power of microarray analysis is given by the possibility of exploring an enormous number of features, which increases the possibility of finding a group that is significant for the question addressed.

This consideration also introduces another commonly discussed point: why measuring RNA and not proteins? One good reason is that proteomic analysis reveals a limited number of proteins (the most abundant) and require large amounts of tissue. Protein aficionados would say that the mRNA level that can be measured in a cell has little to do with the final protein product and that many posttranslational steps influence significantly the activity of a protein, in a way that the mRNA level and/or structure cannot predict. These considerations are welcomed when expression profiling is made to identify a prevalently expressed gene or genes that can be targeted for drug development or for diagnostics. However, if what we are looking for is a profile, the question is probably not relevant. Of course, if there is no mRNA, no protein is expected, unless an extraordinarily stable product was formed from mRNA existing prior to the analysis. The presence of an mRNA does not guarantee the protein, either. However, such cases, in which the mRNA is not translated, are quite anecdotal and often limited to very special contexts, as in the case of oocytes. Huge divergences between mRNA and protein level are also limited to special cases, such as structural or particularly abundant proteins. The extremely interesting field of proteomic analysis will integrate, rather than be in opposition, with gene expression profiling.

Another very common discussion is analysis of microdissected cancers vs whole tumor. Of course, in the case of breast cancer, large variability exists in the nontumoral component, so that the question is perfectly justified. However, many authors argue that the diverse cellular components of a tumor contribute to the tumor biology and should be evaluated as a whole. Indeed, in many studies carried out using nonmicrodissected tumors, genes belonging to lymphocytes, macrophages, adipocytes, stromal cells, and other cell types are clearly visible (16,22). Apparently, however, this does

not mask or hamper the identification of gene expression signatures relevant for clinical associations.

The modern success of RNA analysis on DNA microarrays is eventually linked to its relative ease, reproducibility, robustness, and feasibility and to the amount of information obtained, as compared to other approaches available to date. Once operators become acquainted with RNA extraction. handling, and storage, in fact, probe labeling, hybridization, and data analysis are sufficiently robust and produce satisfactory results most of the time. Analysis of RNA also presents the enormous advantage provided by nucleic acid biochemistry, that is, the possibility of using nanoquantities of material, given that amplification, (i.e., replication) of nucleic acid sequences is always possible with high fidelity. The linear RNA amplification technique allows microarray analysis in a number of relevant applications, notably on laser-captured tissues and on FNA. Laser-capture microdissection allows not only analysis of purified tissue or tumor components, but also comparison of different normal and tumor cells in the same biopsy, for example, different pathological stages concurrently present in the same tumor (normal, hyperplastic, in situ, invasive) (12). Fine-needle biopsies, instead, allow evaluation of gene expression profiling before surgery, for example, when a neoadjuvant protocol can be employed, thus permitting a view of gene expression changes during chemotherapy (11, 23, 37).

Of course, before microarray analysis of breast cancer can become a routine practice, many steps have to be performed and many questions addressed. The first important concern, even more than the kind of microchip (probes, genes) on which an analysis is performed, is given by the kind of reference RNA to which comparison is made. In fact, all the studies published to date used different references, either normal mammary tissues (12), a pool of RNA from a series of cell lines (15,16,29), or a pool of all tumor RNAs (22,30). This poses a series of problems both when comparing results from different laboratories and also in absolute terms. For example, pooling mRNA from many different cell lines can dilute out to very low levels some less abundant, cell-specific mRNA species, thus resulting in abnormal enhancement of the sample-to-reference ratio in some cases, or to unsignificant (and therefore excluded) fluorescence signals in other cases. The debate on the optimal reference RNA is now open and must lead to common reference standards in the near future (38-41). The kind of microarray for analysis is also a question to address. In fact, studies in which different platforms were compared demonstrated a certain degree of discrepancy. Different microarrays use very different probes, for example, 5' vs 3' cDNA fragment, derived from different EST libraries, or oligonucleotide probes directed to different regions of the mRNA, and this may represent a serious problem if genes possess alternative splicing forms. Thus, comparison of microarray studies deserves relevant further work.

It is quite easy to foresee that in the next few years microarrays will expand to cover the entire set of mRNAs encoded by the human genome and that exploration in breast cancer, as well as in cancer in general, will follow the same trend. Initial estimates of the number of genes in our genomes were oriented toward 100,000-150,000, considering the organism complexity and the number of proteins needed. However, the Human Genome Project showed that the actual number of genes is much smaller, currently estimated to be approx 35,000. The reason for this is that vertebrate genomes have evolved alternative splicing to encode subtly different proteins using the same gene. A current estimate is that more than 50% of human genes will encode multiple mRNA and proteins by alternative splicing. It is then likely that future microarrays will accommodate probes for all the splicing forms of all the gens, to allow a complete picture of the encoding capability of our genomes. In parallel, public repositories of data from microarray analysis will be developed, allowing researchers to compare their own data with those of others, and even to mine other investigators' data with new algorithms and data mining tools.

As already exemplified with regard to published studies, genome-wide expression profiling of breast cancer will provide relatively small dedicated sets of genes associated with peculiar characteristics of the tumor. These sets will be useful for prognosis, prediction of response, response monitoring, and prediction of the site of metastasis. It is very likely that biotech companies will develop dedicated microarrays, with hundreds—rather than thousands—of gene probes, to answer specific clinical questions and to perform complete pathological characterization of breast tumors.

As always during transfer of biological knowledge to medicine and industry, extensive technical advances and support implementations are needed, as well as brainstorming among the operators in connection with the new algorithms, bioinformatics, and development of new analytical tools.

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III ASSAYS FOR DNA ABNORMALITIES

5

Assays for Gene Amplification

State of the Art, Pitfalls, and Promises

Chungyeul Kim, MD, Yongkuk Song, PHD, and Soonmyung Paik, MD

CONTENTS

IMPORTANCE OF GENE AMPLIFICATION IN BREAST CANCER HER2 GENE AMPLIFICATION IN BREAST CANCER WHAT IS THE BEST SOURCE OF ASSAY MATERIALS FOR HER2? WHICH IS THE BEST ASSAY PLATFORM FOR THE DETERMINATION OF HER2 STATUS? **REPRODUCIBILITY OF HER2 ASSAYS**— US COOPERATIVE CLINICAL TRIAL GROUP EXPERIENCE **OPTIMIZATION OF FISH** FOR OLD FORMALIN-FIXED PARAFFIN-EMBEDDED TISSUE SPECIMENS CHROMOGENIC IN SITU HYBRIDIZATION **REAL-TIME PCR** COMPARATIVE GENOMIC HYBRIDIZATION (CGH) **CONCLUSIONS** REFERENCES

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SUMMARY

Since the approval of trastuzumab for the treatment of HER2-positive breast cancer (1), there has been an active debate as to the reproducibility and validity of the FDA-approved assays that detect HER2 abnormalities in the tumor tissue. In this chapter, various clinical assays are compared focusing on practical issues.

Key Words: Gene amplification; trastuzumab; *HER*-2 testing; breast cancer.

1. IMPORTANCE OF GENE AMPLIFICATION IN BREAST CANCER

Recent studies using combination of cDNA array based expression profiling and comparative genomic hybridization have elucidated the role of gene amplification in the transcriptional program of breast cancer. In the study by Pollack et al. (2), copy number alteration and expression levels across 6691 mapped human genes were examined in 44 locally advanced breast cancer and 10 breast cancer cell lines. The data from this study suggest that at least 12% of all the variation in gene expression among the breast cancer is directly attributable to underlying variation in gene copy numbers. The total number of genomic alterations (gains and losses) correlated significantly with high grade (p = 0.008), negative estrogen receptor (ER) (p = 0.04), and p53 mutation (p = 0.0006). Of 117 high-level amplifications (representing 91 different genes), 62% (representing 54 genes) were found to be associated with at least moderately elevated mRNA levels, and 42% (representing 36 different genes) with highly elevated mRNA levels. In a similar effort, Hyman et al. examined the correlation between copy number changes and expression levels in 14 breast cancer cell lines using a cDNA microarray of 13,824 genes (3). They found 44% of highly amplified genes resulting in overexpression and 10.5% of overexpressed genes being amplified.

Together these results suggest a profound role of gene amplification in transcriptional control of gene expression in breast cancer and provide a rationale for pursuing amplified genes as a preferred target for developing therapeutics and diagnostics.

2. HER2 GENE AMPLIFICATION IN BREAST CANCER

HER2 was originally cloned as a gene that is amplified with homology to human epidermal growth factor receptor gene. Since Slamon et al. demonstrated the clinical significance of the *HER2* gene amplification (4), numerous studies largely replicated their original findings using a variety of methods ranging from immunohistochemistry to Southern blotting technique.

It is noteworthy that many genes are included in the HER2 amplicon. These include v-*erbA*/thyroid hormone receptor- α (THRA1), the retinoic acid receptor α (RARA), the MLNs 50, 51, 62 steroidogenic acute regulatory protein related protein (MLN 64/CAB-1), peroxisome proliferatoractivated receptor binding protein (PBP/PARBP/TRAP-220), growth factor receptor-bound protein 7 (GRP7), homebox genes 2 and 7 (HOXB2 and HOXB7), junction plakoglobin (JUP), dopamine and cAMP-regulated phosphoprotein (32 kDa in size) (DARPP-32), thyroid hormone receptor associated protein complex component 100 (TRAP-100), titin cap protein (TCP), CDC2 related protein kinase-7 (CrkRS), Aiolos, gastrin, 17B-hydroxysteriod dehydrogenase type 1 (HSD17B1), and topoisomerase IIa (topoIIa, encoded by TOP2A), all of which have been shown to be coamplified in the portion of the HER2-amplified tumors (5). High-resolution mapping of tumors with HER2 amplification with these gene-specific probes may provide additional insight as to the complex behavior of HER2-positive tumors such as differential response to trastuzumab.

3. WHAT IS THE BEST SOURCE OF ASSAY MATERIALS FOR *HER2*?

Trastuzumab, a humanized monoclonal antibody directed against extracellular domain of HER2, is perhaps the most successful targeted therapy second to tamoxifen (1). Because it is difficult to get tissue from many of the metastatic sites at the time of presentation as an advanced disease, the question arises as to whether one can use an archived paraffin block from the primary index tumor as a surrogate for HER2 status of the metastatic lesion. There are only a handful of studies addressing this question of concordance between primary and metastatic lesions, and most of them compared primary and concurrent lymph node metastases. The only true study of concordance comparing primary index tumor with metachronous distant metastases was reported by Niehans et al. (6). They examined HER2 status by immunohistochemistry in 14 autopsy cases (2-9 yr after primary) and found 100% concordance between primary tumors and metastases. In a survey of 56 matched primary and metastatic sites in the node assessed by Herceptest immunohistochemistry, Masood and Bui observed nearly identical staining results between the two (7). The largest study is by Simon et al. (8). In their study, using tissue microarray constructed from 125 cases of breast cancer with nodal metastasis, only 9 cases produced metastasis with partially or completely discordant HER2 status. Among these only two cases exhibited a complete discordance of HER2 status (i.e., all samples of the metastases were HER2 negative). The remaining seven cases exhibited partial discordance (i.e., at least one of their metastases had both HER2-positive and -negative samples). These data seem to justify the use of any tumor tissue,

regardless of whether they are from primary index tumor, metastatic lymph node, or other metastatic site, as a source of material for *HER2* testing. However, owing to the paucity of data for comparing the *HER2* status of index tumor vs systemic metastases, more extensive survey using fluorescence *in situ* hybridization (FISH) is required. Furthermore, if use of trastuzumab in the adjuvant setting is approved, it will be important to examine what happens to the *HER2* status of recurrent tumor cells.

4. WHICH IS THE BEST ASSAY PLATFORM FOR THE DETERMINATION OF *HER2* STATUS?

Because overexpression of *HER2* is almost always attributable to gene amplification (2), one could use assays for protein expression and gene amplification to cross-validate the assays. Thus, there are many available options for the determination of *HER2* status, including immunohistochemistry (IHC), enzyme-linked immunosorbent assay (ELISA), FISH, chromagenic *in situ* hybridization (CISH), and quantitative polymerase chain reaction (qPCR). Each assay has its own advantages as well as disadvantages. The availability of so many options obviously has created confusion among clinicians who are not familiar with the biology of *HER2*.

The most widely used assay for clinical decision making is FISH. FISH is a method in which a fluorescence-labeled DNA probe is directly hybridized to the tissue section and hybridization signals are numerically recorded. In normal cells, because of the presence of two alleles, one each from each parent, one expects to see two distinct signals, whereas in tumor cells with amplification of the target gene numerous signals are seen. In the most popular clinical FISH assay for *HER2* developed by Vysis (PathVysion *HER2* assay), an alpha satellite probe for chromosome 17 labeled with a different color is used to rule out polysomy of chromosome 17 in which the *HER2* gene resides (Fig. 1).

There are two aspects to be considered in making a choice for which assay is to be used for clinical studies. First, one has to consider which test provides better predictability for trastuzumab response. Obviously none of the available assays is ideal, as the response rate of FISH-positive cases to first-line trastuzumab monotherapy is below 50% (9). Dissection of the *HER2* amplicon and signaling pathways may eventually provide better predictors. However, currently, one must use non-ideal tests investigating *HER2* itself. Available data regarding which clinical *HER2* assays are better correlated with clinical response to trastuzumab therapy are not very clear. Much of the confusion is caused by the fact that the IHC assay used in comparison of IHC vs FISH for clinical efficacy in the analysis of pivotal trial data was the Clinical Trials Assay (IHC with both CB-11 and 4D5 antibodies), which is no longer available. Because concordance between Herceptest, an FDA-approved IHC kit, and the CTA was only about 78%,

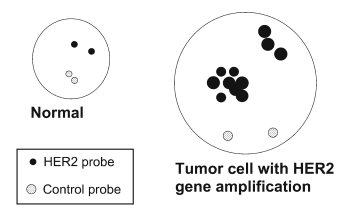


Fig. 1. FISH using Vysis PathVysion assay.

Table 1
H0649g Overall Response Rate
to Single-Agent Trastuzumab (Herceptin)
According to CTA (IHC) or FISH Results

	<i>CTA 3</i> + (%)	CTA 2+ (%)
FISH (+)	22	11
FISH (-)	0	0

Table 2
H0648g Overall Response Rate
to Chemotherapy vs Trastuzumab + Chemotherapy (T+C)

	$CTA \ 3+T+C(\%)$	CTA 3+ Chemotherapy (%)
FISH (+)	55	28
FISH (–)	62	55

it is difficult to extrapolate the pivotal trial data to Herceptest or other IHC assays. However, the data from the pivotal trial for trastuzumab strongly suggest that in the absence of gene amplification as measured by FISH, the benefit from trastuzumab therapy is limited or nonexistent (Tables 1 and 2) (http://www.fda.gov/ohrms/dockets/ac/01/briefing/3815b1_08_HER2%20FISH.doc). Therefore further studies are necessary to answer this question. There are no published data regarding the correlation between gene dosage and trastuzumab response in those cases with gene amplifica-

tion—although logic dictates that such correlations should exist. In summary, the available data suggest that FISH appears to be the assay of choice for predicting response to trastuzumab therapy, but it is clearly not the ideal test.

Other issues to be considered include assay reproducibility, reliability, and practicality. Because most of the treatment decisions for metastatic case are based on analysis of the archived primary index tumor tissue, there is a need for reliable and accurate methods to detect HER2 abnormalities in archived formalin-fixed paraffin-embedded tissue. This requirement essentially eliminates candidates such as ELISA, Northern blot, and Southern blot methods, all of which are better performed on frozen tissue. Although FISH appears to be more accurate and reproducible, complete automation is difficult for FISH. Furthermore, scoring and interpretation of FISH results is time consuming for the pathologists. There is also a consistent assay failure rate of about 5% for FISH. On the other hand, IHC is completely automated and image analysis systems such as ACIS (Chromavision) or ARIOL (Applied Imaging) are now available to help interpret the results. In addition, IHC costs 10-20 times less than the FISH assay. Therefore many laboratories and agencies have now adopted a tiered approach using both IHC and FISH. This is based on the experience by many laboratories that there is a near-perfect correlation between 3+ IHC (by essentially any method using any antibody) and gene amplification by FISH, and between 0 or 1+ IHC and no amplification by FISH. This leaves only 2+ IHC cases that must be confirmed by FISH testing. Dowsett et al. compared the assessment of HER2 by IHC (HercepTest) and FISH in 426 breast carcinomas from patients being considered for trastuzumab therapy (10). The tumors were sent in from 37 hospitals and tested in three reference centers. Only 2/270 (0.7%) IHC 0/1+ tumors were FISH positive. Six of 102 (5.9%) IHC 3+ tumors were FISH negative. Five of the six had between 1.75 and 2.0 HER2 gene copies per chromosome 17 and the sixth had multiple copies of chromosome 17. Thirteen percent of tumors were IHC 2+ and overall 48% of these were FISH positive, but this proportion varied markedly between the centers. Sixty IHC-stained slides selected to be enriched with 2+ cases were circulated among the three laboratories and scored. In 20 cases there was some discordance in scoring. Consideration of the FISH score in these cases led to concordance in the designation of positivity/negativity in 19 of these 20 cases. These data support an algorithm in which FISH testing is restricted to IHC 2+ tumors in reference centers.

This tiered approach is a quite reasonable compromise owing to much less cost and personnel efforts involved. One caveat for this tiered system is that it will work only when qualified laboratories do initial IHC screenings, as the US clinical trial groups learned through their painful experiences from their clinical trials for trastuzumab.

5. REPRODUCIBILITY OF *HER2* ASSAYS— US COOPERATIVE CLINICAL TRIAL GROUP EXPERIENCE

In the United States, the National Adjuvant Breast and Bowel Project (NASBP) is conducting clinical trial B-31, in which node-positive patients are randomized to standard AC-T regimen or the same regimen plus trastuzumab for 1 yr. In this trial original eligibility was based on 3+ IHC results or amplification by Vysis PathVysion assay, reported by any laboratories in the United States or Canada. Owing to patient safety concerns, a central assay for initial 100 cases enrolled into the trial whose blocks become available was preplanned in the protocol. To prevent any bias in assay performance, a third-party commercial reference laboratory was used to perform central assays using both Dako Herceptest (FDA-approved IHC assay) and Vysis PathVysion assay (FDA-approved FISH assay). Results from 104 cases were ultimately evaluated (11). Surprisingly, 19/104 (18%) of the cases were negative by both FISH and IHC (3+) when assayed in the central laboratory. Examination of the data according to where the original assays were performed demonstrated a striking trend between poor concordance and laboratories that handle small assay volumes. Only one of 29 cases from large-volume laboratories (that assay 100 or more cases per month) were incorrectly classified as 3+ by IHC. In contrast, 18 of 75 (24%) cases deemed HER2 positive by small-volume laboratories were negative by both central IHC and FISH. This result was observed regardless of whether FDAapproved Herceptest or other IHC assays were used.

An additional 27 cases were entered based on FISH by local laboratories. When subjected to central assays, all 27 cases were validated to have gene amplification. Seventeen of these cases were entered by small-volume laboratories and 10 from large-volume laboratories. Therefore, we concluded that the FISH assay is in general more reproducible than IHC assays. Based on these findings, protocol eligibility was modified to accept 3+ IHC results only when reported by approved or high-volume laboratories and FISH results by any laboratories. To test the validity of this modification, we conducted a central review of cases entered after the implementation of the eligibility changes. The results confirm our previous findings. Of 240 cases entered after the amendment, only 6 of 133 cases entered based on gene amplification by FISH performed by local laboratories were negative by central FISH. In 1 of 133 cases, a hybridization signal could not be obtained. Among 107 cases that were entered based on 3+ IHC results reported by approved laboratories, 2 were negative by central FISH, with assay failure in 3 cases. Among 6 cases entered by FISH and negative on central assay, 3 had an original *HER2*/Control probe ratio of under 3, suggesting caution when dealing with cases with a ratio under 3. Although our experience

suggests reliability of FISH assay in general, the experience of the Intergroup in its trastuzumab trial is completely different (12). Tumor specimens from the first 119 patients enrolled in its protocol N9831 were centrally tested; 74% were found to be HercepTest 3+ and 66% were found to have *HER2* gene amplification. In contrast to NSABP experience, only 6 of 9 (67%) of the specimens submitted by local laboratories as FISH positive could be confirmed by central assays.

6. OPTIMIZATION OF FISH FOR OLD FORMALIN-FIXED PARAFFIN-EMBEDDED TISSUE SPECIMENS

Although commercial FISH probes are very expensive, one can use DenHyb (Insitus Biotechnologies, Albuquerque, NM) to dilute the probe by approx 1:10 and still obtain comparable results. We have found that this hybridization mix is extremely cost-effective when commercial probes are used (we use 1:15 dilution of the probe mix in Den Hyb for PathVysion kit).

Because *HER2* testing is usually performed using the original primary tumor rather than a metastatic specimen, archived paraffin blocks are often used as test material. In our experience, older blocks have a significant higher failure rate even when we used the Vysis PathVysion *HER2* FISH assay per manufacturer's instructions. However, we have found three generic ways of improving the signal, especially from archived tissue. One is to include a formalin postfixation step during pretreatment before hybridization—this may work especially well for cases with poor fixation. The second method is to increase the incubation time in sodium thiocyanate solution. We generally use longer incubation according to the age of the block with up to 1 h for 30-yr-old blocks. A third method is to use antigen retrieval in pH 4.0 sodium citrate buffer.

In this regard, Anderson et al. have reported a new pretreatment procedure to increase the sensitivity of the FISH assays, with a decrease in assay failure rate from 20% to 10%. Although this protocol is much more complicated than the manufacturer's protocol, it improves the hybridization of older samples (13). Our in-house optimized protocol for Vysis PathVysion *HER2* assay is provided in Table 3.

7. CHROMOGENIC IN SITU HYBRIDIZATION

FISH suffers from two main problems. First, it requires fluorescence microscopy (with special filter sets if using PathVysion) and digital photomicrography for clinical reporting and archiving. Second, pathologists need to identify the area of interest (e.g., delineating the invasive vs *in situ* component), which can be time consuming. Therefore, development of an *in situ*

		NSABP Protocols	NSABP Protocols (accrual period)	
	B-31	B-28	B-14 (1982–1988) and B-20	B-04
Steps	(2000–present)	(1995–1998)	(1988–1993)	(1971–1974)
Deparaffination	3X xylene for 5 min, 2X 100% EtOH for 1 min at RT	Identical	Identical	Identical
Acid treatment	0.2 N HCl for 10 min, dH ₂ O for 3 min	0.2 N HCl for 20 min, dH ₂ O for 3 min	0.2 <i>N</i> HCl for 30 min, dH ₂ O for 3 min	0.2 <i>N</i> HCl for 40 min, dH ₂ O for 3 min
Pretreatment	Vysis pretreatment solution at 80°C for 20 min, dH ₂ O for 3 min	Vysis pretreatment solution at 80°C for 40 min, dH ₂ O for 3 min	Vysis pretreatment solution at 80°C for 60 min, dH ₂ O for 3 min	Vysis pretreatment solution at 80°C for 60 min, dH ₂ O for 3 min
Protease treatment	Vysis Protease I solution at 37°C for 5 min, dH ₂ O for 3 min	Vysis Protease I solution at 37°C for 15 min, dH ₂ O for 3 min	Vysis Protease I solution at 37°C for 20 min, dH ₂ O for 3 min	Vysis Protease I solution at 37°C for 30 min, dH ₂ O for 3 min
Fixation	10% buffered formalin for 10 min at RT, dH ₂ O for 3 min	Skip	Identical	Identical
Dehydration	EtOH series (70, 85, 100%) Identical for 1 min at RT, air dry	Identical	Identical	Identical

hybridization method that questions a colorimetric signal may provide an alternative that would be favored by pathologists.

Tanner et al. have reported a CISH protocol based on hybridization of tissue target DNA with a subtracted fluorescein-labeled probe. Signal amplification is detected using an antifluorescein antibody, and a colorimetric reaction results in brown diaminobenzidine precipitation (14). Subtracted probe technology used in this protocol (subtracting repetitive sequences using subtraction PCR) has the potential advantage over the direct labeled BAC probe used in the PathVysion FISH kit of having less background owing to elimination of repetitive sequences. Furthermore, this method is less expensive, as it employs a PCR amplifiable template for probe generation. CISH produces results very comparable to those obtained with FISH, with 95-100% concordance rate. In our study of 81 cases, identical results for both methods were found in 26 cases (10 amplified, 16 nonamplified) (15). One case was misinterpreted as overexpressed by CISH because of background precipitate. In 4 cases, CISH suggested low-level amplification. Three of these cases subsequently were found to have chromosome 17 polysomy. Therefore, cases with chromosome 17 polysomy and those with background staining may be misinterpreted as amplification signals. However, the clinical consequences of misclassification of polysomy cases are unknown.

Using material from the NSABP B31 trial, we observed hybridization signal from only 60/81 cases. Forty-nine were scored positive for gene amplification by CISH and 11 negative. Among 49 positive cases, 47 were positive by FISH and 2 negative. Among 11 cases negative by CISH, 8 were also negative by FISH and 3 were positive. The overall concordance rate was 92%. In another cohort of 123 evaluable cases from NSABP trial B-15, 88 cases were positive by CISH. Among these, 83 were also positive by FISH. Among 35 cases negative by CISH, 30 were also negative by FISH. Again the concordance rate was 92%. This concordance rate of <100% could simply reflect the fact that we have not fully optimized the assay for the multicenter-derived materials in our trials. Therefore, other published data may be more relevant to general use of CISH.

8. REAL-TIME PCR

Real-time reverse transcriptase-PCR (RT-PCR) permits quantification of DNA. In theory, this method might combine the advantages of nucleic acid and protein measurement. Milson et al. reported results of analysis of 336 cases with real-time PCR method to quantify the copy number changes of the *HER2* gene over a reference gene (β -globulin) (*16*). Real-time PCR gave *HER2/neu* gene doses of 10 for SKBR3 cells and 2 for T47D cells with coefficients of variation (CV) of <3% for within-run and <6% for betweenrun analysis. Examination of 97 breast tumors found a correlation of

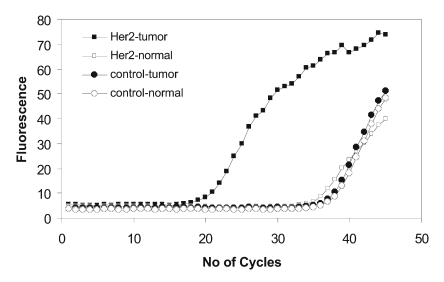


Fig. 2. An example of real-time PCR profile of *HER2* vs control gene in breast cancer specimen with *HER2* amplification.

r = 0.974 between the real-time PCR and quantitative PCR methods. IHC and PCR results agreed for 234 of the subsequent 294 samples analyzed (79% concordance). A subset of 10 discrepant samples was microdissected. After microdissection, all 10 were positive by PCR, thus resolving the discrepancy. We have developed real-time PCR methods based on both TaqMan probes and Cybergreen dye and both work well for formalin-fixed paraffin-embedded tissue. Figure 2 shows representative results from a breast cancer with HER2 gene amplification. One of the major drawbacks of the real-time PCR method is the compression of the dynamic range in comparison to FISH. Only two cycle differences may be observed, even in the presence of a 5- to 10-fold difference in gene copy number. Because of this problem, the assay has to be performed with high precision and quality control. However, owing to the possibility of complete automation using high-throughput devices for PCR, good reproducibility and portability, and low cost per specimen, real-time PCR does have a good potential for clinical assay. Survey of other amplicons also would be possible with relative ease using this approach.

9. COMPARATIVE GENOMIC HYBRIDIZATION (CGH)

Although probably not a good candidate to become a clinical assay, CGH does provide valuable information regarding the status of genome-wide changes in *HER2*-amplified breast cancer. Isola et al compared conven-

tional CGH with FISH and IHC in the same cases (17). The concurrence for *erb*-B2 detection between FISH and IHC was 90%; between FISH and CGH it was 82%, and between IHC and CGH it was 84%. An increased number of losses of 18q and gains of 20q was found in *erb*-B2-positive tumors. *erb*-B2-amplified tumors as detected by FISH, IHC, or CGH had twice as many CGH-defined chromosomal alterations (means of 11.8, 11.0, and 12.7, respectively) as the nonamplified tumors (means of 6.8, 7.0, and 5.6, respectively). Array CGH provides higher resolution than conventional CGH, but in our hands it suffers from the compressed dynamic range when compared to FISH. Therefore actual copy number determination may be difficult.

10. CONCLUSIONS

Currently, the tiered approach for using FISH only for IHC 2+ cases seems to be well justified as long as IHC is performed by experienced laboratories with proper quality control measures. CISH is a promising alternative to FISH. Further dissection of the *HER2* pathway needs to be pursued to develop more sensitive predictors of response to trastuzumab. Investigation of the clinical significance of amplicons other than *HER2* seems to be warranted, based on their contribution to transcriptional control of genes in breast cancer cells.

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IV TISSUE PREDICTIVE BIOMAKERS

Cell Kinetics

Maria Grazia Daidone, PhD, Rosella Silvestrini, PhD, and Dino Amadori, MD

CONTENTS

INTRODUCTION MEASUREMENT OF CELL PROLIFERATION CONCLUSIONS REFERENCES

SUMMARY

Cell proliferative activity represents one of the biological processes most widely investigated because of its association with tumor progression, and in the past years many laboratories have set up and compared different approaches to measure the proliferation of tumor cells for clinical use. Although available results suggest that the majority of proliferation indices may help clinicians in treatment decision making, their clinical usefulness is still controversial owing to some unresolved technical issues linked to preanalytical and analytical aspects and, most importantly, to interpretation of results. However, some laboratories have dedicated considerable time and effort to develop and optimize reproducible methods and standardized methodologies to quantify cell proliferation in clinical tumors, to assess laboratory performance and reproducibility, and to validate preliminary results. Prospective randomized clinical studies have demonstrated the prognostic and predictive significance of breast cancer proliferative activity in different clinical situations.

Cancer Drug Discovery and Development: Biomarkers in Breast Cancer: Molecular Diagnostics for Predicting and Monitoring Therapeutic Effect Edited by: G. Gasparini and D. F. Hayes © Humana Press Inc., Totowa, NJ Novel prospective, multicenter, randomized clinical trials of adjuvant chemotherapy are ongoing to test the utility of cell kinetics to define therapy options for patients with negative or one to three positive nodes presenting rapidly proliferating tumors considered at high risk of relapse. The results will probably help to better establish the predictive role and clinical usefulness of proliferation indices for their transferral to general oncology practice.

Key Words: Assay standardization; cell kinetic-based clinical trials; posttreatment variations; prognostic and predictive relevance.

1. INTRODUCTION

Cell proliferation represents a fundamental biological process because of its involvement in determining growth and in maintaining homeostasis of tissues. The proliferative activity of a tissue can be considered as the result of a complex and dynamic equilibrium of its cell subpopulations. Cells progress through four consecutive phases of the cell cycle, $G_1 \rightarrow S \rightarrow G_2 \rightarrow$ M, under the control of regulatory elements. Cytoplasmic proteins, organelles, and RNA are synthesized in the G₁ and G₂ phases, DNA is replicated during the S phase, and then cells either undergo the mitotic (M) phase or leave the cell cycle and enter a state of quiescence (G_0) . The activity of cell cycle regulators is modulated according to stimulatory or inhibitory growth signals and is subject to strict control in normal cells, whereas in cancer cells a variable degree of independence from such stimuli seems to occur. In particular, the activation of oncogenes, probably associated with the inactivation of tumor suppressor genes, is responsible for the induction of stimulatory signals as well as for the disruption of checkpoints that ensure an orderly progression through the cell cycle (1,2).

The proliferative activity of the tumor cell population, which is closely linked to disease progression, has emerged from clinically oriented research as an increasingly important feature to complement clinicopathological staging for a more accurate prediction of risk of relapse (in different clinical and pathological situations), and has contributed to defining the phenotype of highly aggressive tumors. Thus, in view of biological intertumor heterogeneity, increasing efforts have been made in the past decade to obtain as much cell kinetic information as possible from individual clinical lesions to improve knowledge of tumor biology and potential aggressiveness and to provide clinicians with information on the clinical utility of proliferation markers assessed in large consecutive case series.

To render cell kinetic determination feasible in consecutive series of patients, investigators have turned their attention to specific aspects of the complex phenomenon of proliferation and growth. In particular, kinetic characterization has focused on some specific proliferating cell compartments, that is, on cells that transit through the cell cycle that are generally responsible for tumor growth, and more susceptible to the action of therapeutic agents.

Biologists and pathologists have used several approaches to determine cell proliferative fraction, in accordance with their professional backgrounds (3). Such approaches are based on different rationales and employ different morphometric, immunocytochemical, cytometric, or autoradiographic methods, with inherent advantages and disadvantages. They are designed to analyze and quantify either the whole proliferating fraction or discrete fractions of cells in specific cell cycle phases, mainly in the S and M phases. From a methodological viewpoint, common requirements for clinically useful proliferation markers include technical and biological effectiveness in terms of ability to describe a specific biological phenomenon and to provide results that are informative, reliable, accurate, and reproducible in intra- and interlaboratory settings, at an acceptable cost and obtainable easily and quickly when needed for clinical decision making.

In breast cancer, retrospective correlative studies have indicated that proliferative activity is generally unrelated to clinicopathological stage (4), but at the same time strongly and persistently indicative of risk of relapse and death (5), even in the presence of prognostic information provided by other important pathobiological features. Evidence is also emerging of an association between cell proliferation and response to systemic treatments. Preliminary investigations on the clinical usefulness of cell proliferation have been carried out retrospectively on large, generally monoinstitutional case series, and quality control programs have been activated to guarantee the reproducibility of laboratory determinations (6). Following these initial developments, prospective randomized controlled trials were recently planned to assess the clinical usefulness of proliferation markers, that is, whether they provide information that will affect choice of treatment and improve clinical outcome (7,8). Available results indicate a chemotherapy benefit for node-negative breast cancer patients at high risk of relapse as they present with rapidly proliferating tumors, but the relationship between cell proliferation and response to specific treatments is in need of further validation in large, prospective, proliferation-based studies.

2. MEASUREMENT OF CELL PROLIFERATION

Initially, measurement of the proliferative activity of human tumors was considered with scepticism because it took into account only a fraction of the entire tumor cell population in a single tissue sample obtained at only one time in tumor life, which generally corresponded to surgical intervention for diagnosis and/or tumor removal. Today, currently used proliferation indices assessed by morphometry, immunocytochemistry, flow cytometry, or incorporation techniques have become widely accepted to determine and quantify the whole proliferative fraction or discrete fractions of cells in specific cell cycle phases on consecutive series of clinical tumors, including breast cancers (3,9).

2.1. Phase-Specific Markers

These include proliferation markers measuring the only two phases of the cell cycle in which cells are detectable on the basis of morphological or phenomenological aspects or as a result of their capacity to incorporate DNA precursors.

2.1.1. MITOSIS

Quantification of cells in the mitotic phase (10) is currently expressed as the number of mitotic figures per 10 high-power fields (mitotic activity index [MAI]) or, when corrected for field size and area fraction of the neoplastic epithelium, as standardized mitotic index (volume fraction-corrected mitotic index, or M/VV index, giving the result in mitotic figures per mm^2 of neoplastic epithelium). Both methods of expressing the presence of mitotic figures provide comparable results, but the standardized mitotic index (SMI) has consistently shown smaller interobserver variations. These indices, which have long been employed as diagnostic and prognostic tools in the study of tumor pathology, are important components of all histological grading systems and are routinely used by pathologists. They do not require special processing or staining procedures or the fragmentation of tumor tissues. However, although an increased mitotic activity is a frequent finding in aggressive tumors, the validity of these measurements as markers of tumor proliferative activity will remain controversial (11) until they are standardized or until interlaboratory reproducibility is guaranteed. In fact, mitotic figure counting represents a simple, rapid, and highly feasible approach even for very small tumors, which, however, can be affected by biological and technical factors, and by intra- and interobserver variability owing to the subjective identification of mitotic figures. Although the latter weaknesses can be virtually eliminated by providing precise descriptive criteria for the morphological identification of mitoses, such as those developed by the Amsterdam group (12), technical aspects, including type and time of fixative, section thickness as well as drawbacks related to definition of high-power fields and total number of tumor cells can compromise the interstudy comparability of results. Finally, in addition to problems related to intratumoral heterogeneity and to the poor resolution of the cell kinetics parameter caused by the relatively short time of the M phase (40–60 min) compared to the duration of the entire cell cycle (40-50 h), metaphase arrest may also represent a final stage of cell life.

2.1.2. S-Phase

The quantitative determination of cells in the S phase initially based on the active incorporation of labeled ([³H]thymidine) or halogenated DNA precursors (bromo- or iododeoxyuridine) (13) was successively paralleled by flow or image cytometry of cells with an S-phase DNA content (14). Incorporation measurements, which are performed with autoradiographic or immunohistochemical techniques, require fresh material, aspirates, and surgical or bioptic specimens, and must therefore be prospectively planned. The fraction of S-phase cells is quantified and expressed as the percentage of DNA precursor-incorporating cells over the total number of tumor cells. The main advantages of these approaches, which are considered complex, are the high accessibility and, as *in situ* procedures, the possibility to discriminate tumor from nontumor cells to overcome bias related to tumor heterogeneity. Thymidine labeling index (TLI) is not affected by type or time of fixation, gives clear-cut and unequivocally positive images of reduced silver grains, and permits determinations of labeled cells even after lengthy preservation of archival paraffin blocks.

The main limitation of these approaches is the requirement of fresh tumor material with a sufficient number of viable cells, which has been partially overcome by the availability of kits for TLI (distributed by Euroframe, Asti, Italy) and for bromodeoxy uridine (distributed by Amersham) labeling index (BrdULI) determination, which guarantee the standardization of the first methodological steps and facilitate their use in institutions without adequately equipped laboratories.

The cytometric quantification of nuclear DNA content, which generally provides information on total DNA content and gross genomic abnormalities, can be used to quantify cells in the different cell cycle phases, in particular in the S phase, based on the knowledge that S-phase cells have a variable DNA content ranging from the presynthetic phase $G_{0/1}(2n)$ to the postsynthetic G_2 phase (4n). The utilization of dyes that specifically bind DNA, such as propidium iodide, ethidium bromide, mitramycin, 4',6-diamidino-2phenylindole (DAPI), acridine orange, and Hoechst 33258 allows a quantitation of nuclear DNA content by flow cytometry on isolated nuclei or cell suspensions, or by image static cytometry on cytohistological specimens. Both approaches give a frequency histogram of DNA content, which reflects the cell cycle. The fractions of cells in the different phases are quantified by computerized cell cycle analysis. In addition to S-phase cells, the fraction of cells in the $S+G_2M$ phases is also considered by some authors as a more complete proliferation index that defines the proportion of cells in the cell cycle excluding only those in the $G_{0/1}$ phase. The most diffuse approach for the evaluation of the S-phase cell fraction is flow cytometry (FCM-SPF), the main advantage of which consists in a rapid, potentially objective evaluation of a large number of cells obtained from surgical specimens, biopsy or fine-needle aspirates, effusions, and bone marrow aspirates. The main drawback, which is common to all the non-*in situ* techniques, is the impossibility to discriminate tumor from nontumor cells. This automated technique received a major impetus in the late 1980s, with the development of procedures to perform flow cytometry in solid tumors using material from formalin-fixed paraffin-embedded blocks (14) or from frozen tumor specimens. The use of the latter material also guarantees more reproducible information on specimens that have been in storage for some time and accrued from different centers. The feasibility of FCM-SPF is potentially high, but the quality of results can be affected by methodological factors. To make results reproducible and comparable among the different centers, standardization of assay methodologies, cell cycle analysis techniques, and cutoff points for classifying and interpreting FCM-S from DNA histograms, as well as strict quality control, are mandatory.

Recently, a concerted effort was carried out and developed by US, French, and Swedish investigators to optimize the prognostic strength of flowcytometric DNA measurements and to test the validity of the proposed adjustments (15). This study, performed on about 1400 patients with nodenegative breast cancer, emphasized the complexity of the interpretation of DNA ploidy histograms and quantification of S-phase cells, which are so closely related to each other that they provide non-independent prognostic information when considered in association. Following 10 adjustments to the two measurements, which involved both DNA ploidy reclassification and S-phase calculation, the association between the two flow cytometric measurements has been reduced and their confounding technical correlation eliminated, thus permitting them to become independent prognostic factors in a single model.

2.2. Antigen-Related Markers

These approaches involve the determination of enzymes (DNA polymerase α , thymidine kinase), antigens (Ki67/MIB-1, KiS1, KiS2, cyclin-PCNA), or structural alterations (AgNOR, that is, argyrophilic nucleolar organizer regions) (16–21), which in some instances represent the natural evolution and the integration of morphological and functional determinations, and provide information on the overall fraction of proliferating cells, that is, the growth fraction of the tumor. However, available information on their clinical relevance, albeit interesting, has been questioned, especially for solid tumors, and indicates the need for methodological verification and standardization through quality control assessments (10).

Among proliferation-associated antigens, Ki67 has long been regarded as the most reliable marker of proliferating cells. The expression of Ki67 antigen, detectable by Ki67 antibody, was first identified in phytohemagglutinin-stimulated lymphocytes and described as putatively expressed only by proliferating cells (18). Its main drawback is that it can be detected only in acetone-fixed frozen sections. However, the recent availability of a series of reagents (MIB-1, Ki-S1, and Ki-S5), which recognize Ki67 proliferation antigen and can be used on formalin-fixed paraffin-embedded material following antigen retrieval with pretreatment in a microwave oven or in a pressure cooker (10), has overcome the fixation-related constraint and opened up interesting perspectives for assessing the clinical utility of MIB-1 index on consecutive large series of cancers.

2.3. Comparability and Standardization of the Different Proliferation Indices

The indices most frequently used to quantify the proliferative rate of clinical tumors (mitotic figures, FCM-SPF, TLI, and Ki67/MIB-1) can be determined on viable, frozen, or paraffin-embedded tissues using different detection techniques, which may also involve DNA histogram modeling software. Each approach has inherent advantages and disadvantages, including different feasibility rates, which appear to depend on the availability of fresh tumor tissue for TLI or BrdULI, and on data analysis and interpretation in tumors with multiple cell subpopulations for FCM-SPF. These differences in biological and technical aspects could be one of the main causes of the inconsistency of results among laboratories, which may help to explain the reasoning behind the caution raised by the American Society of Clinical Oncology against the routine use of proliferation indices in breast cancer, notwithstanding the large number of studies demonstrating the prognostic relevance of the different tumor proliferation indices. Prospective studies, associated with an evaluation of laboratory effectiveness in terms of presence of quality assurance programs and methods comparison, could provide definitive information on the clinical utility of proliferation markers.

2.3.1. Reproducibility of the Measurements and Quality Control Programs

Efforts have been made to standardize methodologies and interpretation criteria to improve reliability, accuracy, and reproducibility of results within and among the different laboratories and to promote and maintain quality control programs to provide clinicians with a network of qualified laboratories for currently employed proliferation indices. In particular, it is worth mentioning that all the prospective randomized phase III trials of chemotherapy vs observation in node-negative breast cancer, activated in the last decade to test the clinical utility of identifying high-risk patients on the basis of high tumor cell proliferation (by MAI, TLI, or FCM-SPF), have been paralleled by the activation of quality control programs for preanalytical and analytical phases of cell kinetic determination (6, 12, 22-25).

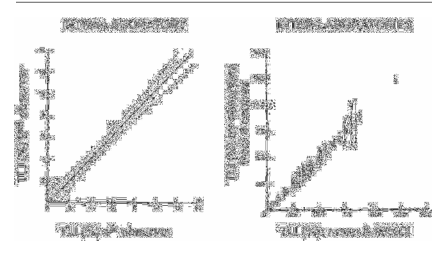


Fig. 1. Reproducibility of TLI determination within a single laboratory (intralaboratory) and among eight different laboratories (interlaboratory) participating in the Italian Network for Quality Assessment of Tumor Biomarkers (6).

The external quality control assessment of TLI determination has been extended nationwide in Italy (22) to ensure its reproducibility, and is currently used in multicenter clinical protocols. This initiative focuses on the assessment of the reproducibility of histoautoradiographic evaluation of samples and, in accordance with previously defined standard operating procedures for sample scoring, is based on (1) sample identification and selection by the coordinating laboratory; (2) circulation of samples among the participating laboratories; (3) TLI evaluation; and (4) data collection and analysis by an independent Statistical Unit. Results (6) to date have demonstrated satisfactory levels of intra- and interlaboratory reproducibility (Fig. 1), with an improvement in the performance of participating laboratories to more than three runs.

2.3.3. Association Between Proliferation Indices: Basic Results and Clinical Studies

The different indices of proliferation have not always proven to be associated with one another in terms of biological or clinical significance when comparatively analyzed in the same case series. In fact (Table 1), moderate or poor correlation coefficients have generally been observed between proliferation indices detecting cells not only in different but also in the same cell cycle phase, and slightly varying sensitivity and specificity rates have been reported for different proliferation indices when related to clinical outcome on the same case series (21, 26-49). However, at present only one study (29)has been specifically planned to simultaneously challenge the prognostic

	Comparison	Betw	een Different F	Prolifera	tion Markers	
	SPF		Ki67/MIB	-1	MI-N	1/V
	No. of cases (No. of studies)	r	No. of cases (No. of studies)	r	No. of cases (No. of studies)	r
TLI/BrdULI SPF Ki67/MIB-1	1078 (6)	0.54	1482 (7) 1835 (23)	0.29 0.42	541 (2) 611 (3) 1046 (7)	0.27–0.81 0.42 0.49

Table 1	
Comparison Between Different Proliferation Mar	kers

MI, Mitotic (figure) index/count; M/V, volume/corrected mitotic index; r, median correlation or regression coefficients.

Space constraints do not permit us to mention all the articles whose results contributed to the data reported in the table.

capability of BrdULI, MI, or MIB-1 on the same series of stage I-III breast cancers, and its results support the importance of MI evaluation over the other proliferation markers.

2.3.3.1. Association With Risk Profiles

Initial studies investigated the association between cell kinetic markers and the most important conventional prognostic factors, such as the extent of the disease at diagnosis, that is, at the time of clinical detection. Information is available for TLI on 14,147 primary breast cancers collected and characterized in a single institution, the National Cancer Institute (Istituto Nazionale Tumori) of Milan, over a 25-yr period (1975–2000). Proliferative activity was only weakly related to tumor size, as large tumors (>2 cm) were slightly more frequent in the highest proliferation class, whereas it was unrelated to nodal involvement (Fig. 2). Conversely, in undifferentiated tumors and in premenopausal women or in women younger than 50 yr of age, high cell proliferation was more frequently observed (Fig. 2). However, such associations, although statistically significant, are relatively weak. A direct association was much stronger with biomarkers that are indicative of an unfavorable clinical outcome, such as an uploid DNA content, a weak or absent bcl-2 expression, absence of steroid hormone receptors, and p53 accumulation (Fig. 3). Moreover, when considering the association between proliferation and traditional prognostic factors for breast cancer including tumor size, patient age, histological grade, estrogen and progesterone receptors (ER, PgR), the more unfavorable factors that are detected in a tumor, be it node negative or node positive, the more likely it is to be rapidly proliferating. In fact, when four or more unfavorable factors (i.e., size >2 cm, age <50 yr, ER-negative, PgR-negative, grade 3) were simultaneously present, even in the absence of axillary lymph node involvement, tumor proliferative activity was very high in about 30-40% of cases

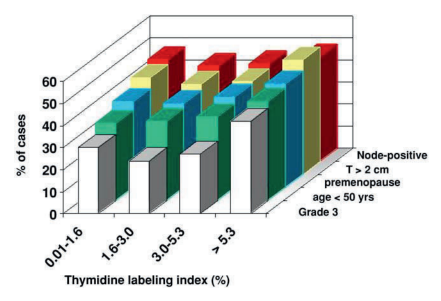


Fig. 2. Unfavorable clinicopathological factors (including patient age and menopausal status, tumor size, axillary lymph node involvement, histological grade) as a function of TLI in 14,147 primary breast cancers.

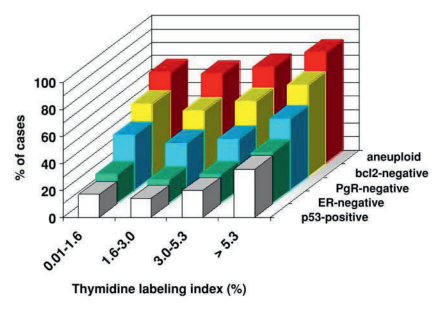


Fig. 3. Unfavorable biological factors (including DNA-ploidy, estrogen and progesterone receptor [ER and PgR] status, bcl-2 and p53 expression) as a function of TLI in 14,147 primary breast cancers.

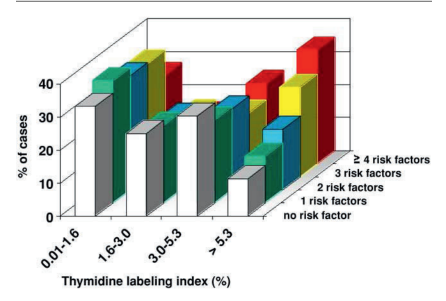


Fig. 4. Relationship between the simultaneous presence of unfavorable clinicopathological and biological factors (including tumor size >2 cm, patient age <50 yr, ER-negative, PgR-negative, histological grade 3) and TLI in nodenegative primary breast cancers.

(Fig. 4). Conversely, the highest proliferation class proved to have only 10-13% of cases with no or only one unfavorable factor, which were more frequent (33 and 36% of cases, respectively) in the lowest proliferation subset.

2.3.3.2. Prognostic Significance

Translational research into human tumors has been developed along two main lines: (1) the search for markers to use as a complement to clinicopathological staging to identify patients at minimal risk of relapse or those who are destined to relapse and progress regardless of treatment and (2) the prediction of patients who are likely to respond or develop resistance to a specific treatment. At present, cell proliferation indices are used to identify patients at high risk of relapse or death (who need aggressive treatments) and those with an indolent disease (who are potentially curable by locoregional treatment alone). In fact, the majority of published articles identified in the PubMed database using *breast cancer* and the name of each *proliferation index* as search terms indicate a direct association between high proliferation indices and the probability of relapse, mainly in distant sites, and death in univariate analysis (Table 2), both in patients subjected to locoregional therapy alone until relapse or in those given adjuvant systemic treatments after radical or conservative surgery. This finding has been

		5	ith worse outco iferating tumo	
	No sys treatr		Different treatm	-
	Relapse	Death	Relapse	Death
TLI, BrdULI				
Univariate analysis	10/10	7/8	8/9	7/7
Multivariate analysis	9/9	7/7	4/4	4/4
FCM-SPF				
Univariate analysis	18/20	8/10	23/26	19/22
Multivariate analysis	10/10	5/5	14/14	13/13
Ki67, MIB-1				
Univariate analysis	4/5	3/3	14/16	10/11
Multivariate analysis	2/2	2/2	6/6	3/3
MI, MAI, M/V				
Univariate analysis	2/3	2/2	15/15	11/11
Multivariate analysis	2/2	1/1	8/8	7/7

Table 2
Prognostic Value of Proliferation Indices in Breast Cancer

^{*a*}Identified by a computerized literature search performed by PubMed using *Breast* cancer and the name of each of the *proliferation indices* as search terms, on all available original English articles that were selected for inclusion when they reported data on the relationship between proliferation indices and clinical outcome, retrospectively evaluated in univariate and/or multivariate analyses on independent case series of at least 100 patients with a minimum follow-up of 4 yr.

BrdULI, Bromodeoxyuridine labeling index; FCM-SPF, flow-cytometric S-phase cell fraction; MAI, mitotic activity index; MI, mitotic index; M/V, mitotic activity (volume/corrected MI); TLI, [³H]thymidine labeling index.

confirmed for all the proliferative indices and regardless of the criteria used to classify tumors as slowly or rapidly proliferating. In all these phase I and II exploratory investigations, carried out over the last two decades without *a priori* study design or prospective definition of specimen collection procedures (studies providing level of evidence [LOE] III according to the Tumor Marker Grading Utility System [50]), such an association was generally maintained in multivariate analyses including patient age and menopausal status, tumor size, regional lymph nodal status and histological/ cytological findings or biological markers associated with differentiation, hormone responsiveness, neo-angiogenesis, and genomic alterations (30–32,35,42,51–80) (Table 3). In particular, phase-specific proliferation indices, including mitotic figure counts, maintained their predictivity for disease-, event-, or relapse-free survival and for overall or cancer-specific

survival even in the presence of information provided by histological or nuclear grade, despite the fact that all grading systems included the proliferative information provided by the mitotic index (*35,69,73,75,77,78*).

From such an overview it also emerged that, in the presence of other clinical and pathobiological factors, proliferation indices contributed to a better prognostic definition within subsets of patients with an intermediate tumor size (1–2 cm) (55), or those classified at an intermediate risk of relapse (5) according to the criteria adopted in the 1998 St. Gallen International Consensus Conference on the Treatment of Primary Breast Cancer (81) and integrating patient age, tumor size, histological grade, and steroid hormone receptors (Fig. 5). These findings were confirmed in a large clinical trial conducted by Intergroup (INT 0102), with the maximum level of evidence (LOE I) as it was a prospectively randomized study, in which a low FCM-SPF identified patients with ER/PgR-positive tumors of ≤ 2 cm who would have a clinical outcome similar to that of women whose tumors were too small for biochemical ER/PgR assay (82), and who could be subjected to locoregional treatment alone.

In keeping with these observations are the findings obtained on a series of 2670 patients with histologically node-negative operable breast cancers who underwent radical mastectomy (38.6% of cases) or conservative surgery plus radiotherapy (61.4% of cases) and axillary lymph node dissection at the National Cancer Institute of Milan from 1975 to 1996, and who did not receive any systemic postoperative therapy until new disease manifestation was documented. In this case series, which was consecutive with respect to TLI determined at the time of diagnosis and in which very small $(\leq 1 \text{ cm})$ and large (>2 cm) tumors accounted for 12.2% and 34.6% of cases, respectively, pathological tumor size was a strong predictor of both 10-yr relapse (size $1-2 \text{ cm vs} \le 1 \text{ cm}$; HR = 1.33, 95% CI, 1.04–1.69; two-sided p = 0.0217; size >2 cm vs ≤ 1 cm; HR = 1.82, 95% CI, 1.42–2.33; two-sided p = 0.0001) and death (size 1-2 cm vs ≤ 1 cm; HR = 2.11, 95% CI, 1.36-3.28; two-sided p = 0.0009; size >2 cm vs ≤ 1 cm; HR = 3.75, 95% CI, 2.42–5.81; two-sided p = 0.0001). In patients with very small tumors, relapse-free survival was predicted only by cell proliferation (high vs low TLI, HR = 2.04, 95% CI, 1.23–3.28; two-sided p = 0.005), even in the presence of information provided by patient age and other tumor biological features, such as steroid hormone receptors or p53 and bcl-2 expression. Similar findings were observed for intermediate size tumors (1-2 cm, accounting for 53.2% of cases) in which, in addition to TLI, bcl-2 expression proved to be associated with prognosis (high vs low TLI, HR = 1.31, 95% CI, 1.03-1.68; two-sided p = 0.03; low vs high bcl-2, HR = 1.36, 95% CI, 1.07–1.75; two-sided p = 0.014). Conversely, when the 10-vr probabilities of disease outcome were plotted against continuous values of TLI, the relationship between relapse-free survival and cell proliferation appeared to be differ-

	V	vith Early B	reast Cancer	Subjected to Or	with Early Breast Cancer Subjected to Only Surgery \pm Radiotherapy Until Relapse		
		No.	Follow-un	Proliferation	Other factors considered	p value	ne
Author (ref.)	Stage	of cases	(yr)	Index	for survival analyses	Relapse	Death
Medri (51)	\mathbf{N}_{-}	378	5	TLI	A, T, HG, ER, PgR, MVD	0.08	
Paradiso (52)	\mathbf{N}^{-}	101	5	TLI	A, T, HG, ER, PgR	0.04	
Silvestrini (53)	\mathbf{N}_{-}	215	5	TLI	A,T, ER	0.004	0.035
Courdi (54)	\mathbf{Z}^{I}	167	8	TLI	A, T, HG, ER, PgR	0.037	
Silvestrini (55)	\mathbf{N}^{-}	1800	8	TLI	A, T, ER, PgR	0.006	0.014
Meyer (30)	\mathbf{N}^{-}	414	10	TLI	A, T, ER, nuclear size	NS	0.095
Cooke (56)	Mixed	164	8	TLI	T, N, HER-2/neu		NS
Tubiana (57)	Mixed	125	15	TLI	T, HG	<0.05	0.05
Aubele $(58)^a$	Ι	329	8	SPF	Morphometric features, ploidy	NS	
Sigurdsson (59)	\mathbf{N}^{-}	250	4	FCM-SPF	A, T, ER, PgR, ploidy	0.03	0.0001
O'Reilly (60)	\mathbf{N}^{-}	169	5	FCM-SPF	T, HG, ER, ploidy	0.05	0.045
Brown (42)	\mathbf{N}^{-}	314	9	FCM-SPF	A, T, ER, PgR, cell type,		
					surgery, treatment, Ki67, ploidy	0.006	
Harbeck (61)	\mathbf{Z}^{-}	125	6.5	FCM-SPF	T, HG, ER, PgR, PAI-1, uPA, cathepsin D,	NS	
					p53, HER-2/neu		
Merkel (62)	\mathbf{Z}^{-}	280	6.5	FCM-SPF	T, HG, NG	NS	0.05
Winchester (32)	\mathbf{Z}^{-}	198	6.5	FCM-SPF	A, T, HG, NG, ER, MAI, ploidy	0.03^{b}	
Baslev (63)	\mathbf{N}^{-}	421	L	FCM-SPF	A, T, HG, ER, PgR, no. of examined N	NS	NS
Stal (64)	\mathbf{Z}^{-}	152	8	FCM-SPF	A, T, ER	0.035	
Isola (65)	\mathbf{Z}^{-}	289	8.5	FCM-SPF	T, HG, ER, ploidy, p53, HER-2/neu		0.0001

 Table 3

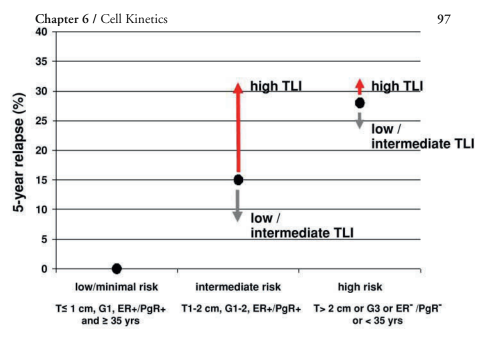
 Prognostic Value of Proliferation Indices by Multivariate Analysis in Patients

	0.03 NS NS	NS	NS		NS	0.003		0.08	0.02			NS		NS	(continued)
NS	0.03	0.05		0.038 0.020		0.022		0.04			0.03	NS)
Peritumoral lymphatic and vessel invasion,	T, HG, ploidy, HER2/neu T, HG, histotype, MI, M/V, ploidy A, T, HG, NG, ER, tubule formation, vascular invasion, necrosis, desmoplasia,	Inflammatory reaction, MI, plotdy A, T, HG, NG, ER, no. of examined N, histotype, ploidy	T, HG, ER, PgR, nuclear pleomorphism, mitotic count	ER, N, clinical stage, ploidy T. HG. N. nathological stage, ploidy	A, T, ER, N, ploidy, HER-2/neu	A, T, HG, ER, PgR, N, histotype,	nuclear pleomorphism, intraductal growth, treatment, M/V, ploidy	A, T, HG, clinical N	A, T, HG, N, histotype, tumor margin,	tubule formation, nuclear pleomorphism, necrosis, intraductal growth, MAI	A, T, HG, ER, histotype	T, HG, tumor necrosis,	lymphatic and blood vessel invasion, p53, HER-2/neu, PCNA	T, HG, histotype, SPF, ploidy	
FCM-SPF	FCM-SPF FCM-SPF SPF	FCM-SPF	FCM-SPF	FCM-SPF FCM-SPF	FCM-SPF	FCM-SPF		FCM-SPF	FCM-SPF		Mitotic count	Mitotic count		Mitotic count	
6	9 10 10	12.5	S	5.5 6	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	6		10	25		6	10		10	
136	100 180 115	265	123	210 158	201	148		377	223		98	200		180	
\mathbf{N}^{-}	$\mathbf{Z} \mathbf{Z} \mathbf{Z}$	\mathbf{N}^{-}	NO	Mixed	Mixed	Mixed		Mixed	Mixed		\mathbf{N}^{-}	\mathbf{N}^{-}		N	
Bosari (66)	Johnson (67) Lipponen (36) Peirò (31) ^a	Witzig (68)	Joensuu (69)	Klintenberg (70) Arnelov (71)	Stanton (72)	Eskelinen (35)		Fisher (73)	Toikkanen (74)		Ladekarl (75)	Kato (76)		Lipponen (36)	

		No.	Follow-up	Follow-up Proliferation	Other factors considered	p value	lue
Author (ref.)	Stage	<i>io</i>	(yr) ¹	Index	for survival analyses	Relapse	Death
Aaltomaa (77)	N	294	13	Mitotic count	A, T, HG, histotype, tumor margin, tubule formation. necrosis.	0.005	NS
Clayton (78)	$^{\rm N}_{\rm -}$	378	20	Mitotic count	intraductal growth A, T, HG, NG, ER, lymphatic invasion, skin/muscle invasion, nucleoli		<0.001
Joensuu (69)	NO	161	5	Mitotic count	T, HG, ER, PgR, nuclear pleomorphism, SPF		<0.0001
Eskelinen (35)	Mixed	216	6	Mitotic count	A, T, HG, ER, PgR, N, histotype, treatment	0.010	NS
Toikkanen (74)	Mixed	223	28	Mitotic count	A, T, NG, N, histotype, tumor margin,		NS
					tubule formation, nuclear pleomorphism,		
Dround (12)	Z	619	9	LY:A	A T ED DED cell true control produced	0.0005	
Iaconetta (70)		263		MIR-1	Timor eize histological grade FR DoP 53	SNC SNC	SN
)		HER-2/neu	2	2
Harbeck (40)	\mathbf{Z}^{-}	100	6.5	MIB-1	T, HG, ER, PgR, PAI-1, uPA, cathepsin D, p53. HER-2/neu	NS	NS
Pinder (80)	Mixed	177	12	MIB-1	T, HG, N	0.05	
A, Patient age; 1	HG, histolc	gical grade;	MVD, microve	essel density; N, a	A, Patient age; HG, histological grade; MVD, microvessel density; N, axillary nodal status; NG, nuclear grade; NS, not significant; T, tumor size.	ot sign	ot significant; T, t

Table 3 (continued)

^{*a*}Image cytometry. ^{*b*}Diploid tumors.



Daidone & Silvestrini, JNCI Monographs, 30, 2001

Fig. 5. Prognostic contribution, in terms of 5-yr relapse probability (%), provided by TLI ($\leq 3\%$, gray line; > 3%, black line) to risk categories of nodenegative breast cancers based on patient age, tumor size, histological grade, and ER and PgR status (80). Dots represent the 5-yr relapse probabilities (%) calculated for risk categories. *Arrows* indicate the 5-yr relapse probabilities (%) calculated as a function of tumor proliferation classes (according to TLI) within each risk category. Overall analysis performed on 543 cases (5).

ently modulated as a function of tumor size (Fig. 6). In fact, a direct inverse relationship between relapse-free survival probability and TLI values was observed in the subset of patients with very small tumors and was also paralleled by the pattern of relationship observed for women with intermediate-size (1–2 cm) tumors, whereas in those with large tumors, relapse-free survival sharply decreased from about 70% to about 45% by raising the TLI cutoff value to 3%, after which this relationship remained almost constant.

2.3.3.3. Predictivity of Response to Specific Treatments

The second objective of translational research in breast cancer, that is, the identification of patients who are likely to respond or develop resistance to a specific treatment, has not provided conclusive results on the role of proliferation indices. The reason for this is probably the generally modest study design, which is not appropriate for investigating the relevance and utility of biomarkers within the context of clinical treatment protocols. In fact, the ideal method to assess the role of any biological variable as a

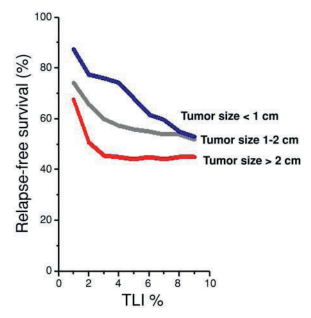


Fig. 6. Relationship between values of TLI and percentage probability of 10-yr relapse-free survival as a function of pathological tumor size in 2670 node-negative operable breast cancer patients not receiving any postoperative systemic adjuvant therapy.

predictor of response to a specific treatment would entail its prospective evaluation in a randomized clinical study specifically designed to analyze biomarker predictivity, or to compare systemic treatments with locoregional therapies. However, to date, only a small number of prospective, high-powered studies addressing the issue of the clinical utility of proliferation markers has provided conclusive results (presented and discussed in Section 2.3.3.4.). Conversely, most of the available results have been retrospectively derived from studies performed mainly in an adjuvant setting, in which the advantage of a long-term follow-up has been counterbalanced by a marked heterogeneity in technical and analytical procedures for biomarker determination on tumor specimens collected for a variety of reasons and available for evaluating proliferation indices without any *a priori* study design.

In advanced lesions, in general, a benefit from intensive polychemotherapy, including S-phase–specific agents, has been observed for rapidly (83-85) rather than slowly proliferating tumors. A greater benefit for patients with rapidly proliferating tumors has also been observed, albeit not consistently, in adjuvant settings for node-positive resectable cancers (86-90). Within the context of prospective randomized-clinical trials, a benefit from cyclophosphamide, methotrexate, and fluorouracil (CMF) on long-term clinical outcome has been observed (88) for both subgroups, although more evidently (87,90) in patients with rapidly proliferating tumors, but also independently of proliferative activity (89). The former finding has been ascribed to a higher efficacy of polychemotherapy, including antimetabolites, in killing dividing cells. In addition to the type of drugs used, treatment schedule has also been proven to affect the relationship between cell kinetics and clinical outcome. In fact, in a randomized treatment protocol aimed to compare the efficacy of alternating vs sequential regimens of doxorubicin and CMF in node-positive breast cancer patients, the benefit of sequential administration was evident mainly in patients with low to intermediate proliferative tumors (91). This result was explained by a partial synchronization of cells in the G_2 –M phases following the administration of high dose intensity doxorubicin and the subsequent presentation of a large fraction of S-phase cells to antimetabolites included in the CMF regimen.

Recently, neoadjuvant chemotherapy protocols have been used as an ideal model for translational studies aimed at analyzing the predictivity of biological variables for short- and long-term clinical end points and at monitoring, at cellular and molecular levels, the effect of treatment by sequential determinations of biomarkers within a single tumor, in the presence of only intralesional heterogeneity (92). Overall, proliferative activity appears to provide information on tumor biological changes and is sensitive enough to reflect, at the cellular level, the biological downstaging induced by treatment, with relevant implications on long-term follow-up. In fact, significant changes observed after chemotherapy generally consisted in a reduction in proliferative activity (93–97). Tumor shrinkage proved to be less frequent in patients with pretreated slowly proliferating tumors (93–99), while a favorable long-term clinical outcome was generally (99) observed for patients with posttreated slowly proliferating tumors (96, 100).

A reduction in the proliferative rate associated with a higher probability of tumor shrinkage and a more favorable outcome for patients with posttreated slowly proliferating tumors was also observed following neoadjuvant antihormonal treatment (101,102). A higher response rate was observed for patients with slowly rather than with rapidly proliferating tumors (103), even in the ER⁺ subset (104,105). These findings, which held true regardless of PgR status, indicate that slowly proliferating metastatic ER⁺ breast cancers benefit from tamoxifen, whereas rapidly proliferating tumors largely escape endocrine control, even if they are ER⁺, that is, traditionally considered hormone responsive. Therefore, in addition to a lack of estrogen receptors, a high proliferative activity represents a limiting factor for response to endocrine therapy. Furthermore, evidence of a major benefit from endocrine therapy for slowly proliferating tumors has also been confirmed in adjuvant settings (106–108), although contrasting results are also present in the literature (109).

However, it must be remembered that these data were obtained mainly from retrospective clinical analyses, with a LOE III (50), and are insufficient to be able to draw definitive conclusions on the predictive role of proliferation indices or to use for therapeutic decision-making (endocrine therapy or chemotherapy). Moreover, they are only suggestive of qualitative relationships, and appropriately designed prospective studies are needed to confirm and validate preliminary results to define whether the improved clinical outcome for patients with ER+, slowly proliferating cancer is the result of a natural indolence or of a specific susceptibility to endocrine treatment, and to evaluate whether the benefit of polychemotherapy with S-phase-specific agents in patients with rapidly proliferating tumors is attributable to a quantitative relationship between drug effect and fraction of sensitive S-phase cells. Recently, the determination of cell proliferation was prospectively planned within the context of adjuvant and neoadjuvant treatment protocols in which the evaluation of the usefulness of biological information represented a secondary objective of the clinical study. Even though such studies are not specifically designed to test the predictivity of proliferation indices, it is likely that they will improve the quality of information on their predictive accuracy and provide a definitive evaluation of the clinical usefulness of a cell kinetic characterization.

2.3.3.4. Clinical Utility

All previous findings demonstrate that the various proliferation indices could provide, with a variable level of specificity and sensitivity, information to identify patients (1) at minimal risk of relapse; (2) destined to relapse and progress regardless of treatment; (3) likely to respond or develop resistance to a specific treatment. However, it is only in the past few years that these results, albeit extremely interesting, have been challenged with respect to the clinical usefulness of cell kinetic information compared to or in association with other pathobiological information. Recently, in fact, the results became available of the first therapeutic clinical trials in which the determination of proliferation indices was planned *a priori*, with a sufficiently powered study design to define the role of prognostic/predictive markers. These provided preliminary information on the actual utility of a cell kinetic determination in the presence of a risk profile defined by other clinicopathological and biological factors.

LOE II studies (50), companion to therapeutic clinical trials, provided evidence in favor of the following:

1. A contribution of cell proliferation evaluated as FCM-SPF to define, in association with patient age, PgR status and tumor size, a broad spectrum of clinicopathobiological categories with a different 10-yr risk of distant metastases within a subset of 800 node-negative ER⁺ cancer patients given tamoxifen (*110*). The risk probability ranged from 70%

(for patients under 35 yr of age with large, PgR-negative tumors and a very high FCM-SPF) to 20% (for patients older than 50 yr of age with PgR-positive, 1-cm tumors and a negligible proliferative activity), and, on an overall 1118 women with node-negative invasive breast cancer up to 5 cm in size, in which Ki67/MIB-1 was considered in addition to FCM-SPF;

- 2. A favorable prognosis for women with slowly proliferating tumors, with superimposable disease-free survival rates for patients who received only surgery or adjuvant tamoxifen or doxorubicin and cyclophosphamide (111);
- 3. A survival advantage for patients with rapidly proliferating tumors who received postoperative adjuvant chemotherapy with doxorubicin and cyclophosphamide, with disease-free survival similar to that of patients with slowly proliferating tumors (112).

Such information provides an accurate assessment of individual patient prognosis and suggests that an aggressive therapy is indicated for only some of the women with node-negative tumors, that is, for those presenting with rapidly proliferating cancers. In keeping with this line of evidence are the findings provided by the prospective randomized clinical trial (LOE I) conducted by the U.S. Intergroup, in which FCM-SPF was able to identify, within the "uncertain" risk subset (ER- or PgR-positive tumors ≤ 2 cm), patients at a low or high risk of relapse (82).

The successive step of prospective translational studies was to investigate whether node-negative breast cancer patients defined at high risk on the basis of tumor cell proliferation could benefit from adjuvant polychemotherapy. One monoinstitutional and two multicenter phase III randomized trials using TLI (7,8) or MAI (113) were activated in Europe, in which the prognostic factor hypothesis was combined with a treatment hypothesis (114). In these studies, patients with node-negative breast cancer were stratified into low- and high-risk groups based on the proliferation index of the primary tumor. Patients with slowly proliferating tumors were not treated with systemic therapy following radical or conservative surgery plus radiotherapy. Patients with rapidly proliferating tumors were randomized to receive or not adjuvant chemotherapy (CMF or FAC). Activation of these trials, as well as of similar studies involving the determination of proliferation indices, was paralleled by the promotion and maintenance of quality control programs for analytical and preanalytical phases of cell kinetic determinations (6.12, 23, 24, 115).

Results are available from the study by Amadori et al. (7) and Paradiso et al. (8), in which 278 and 248 patients with histologically assessed nodenegative tumors were randomized to receive or not to receive chemotherapy. Survival curves showed a disease-free survival benefit in CMF or FACtreated compared to untreated patients (83% vs 72% [7] and 81% vs 69% [8]), with a reduction in both locoregional and distant relapses. It was also observed (7) that the benefit from CMF treatment was evident mainly for cases at very high risk, that is, with the highest TLI values.

Overall, these results support the use of cell proliferation in the nodenegative subset to identify patients at low or minimal risk of relapse and spare them from aggressive treatments, and to select those at high risk as candidates for systemic adjuvant therapy. The finding of a greater benefit from antimetabolite-based regimens in patients with tumors with the highest proliferative rate is in keeping with evidence of a relationship between cell kinetics and response to specific treatments, previously discussed in Subheading 2.3.3.

3. CONCLUSIONS

Proliferation indices can be considered markers of clinical utility. In fact, in node-negative breast cancers the usefulness of the different proliferation indices to identify subsets at very low risk of relapse has been assessed in large retrospective studies and validated in prospective studies (82,111). A benefit from antimetabolite-including chemotherapy regimens has been indicated from retrospective analyses performed in companion studies of randomized treatment protocols and assessed in phase III prospective confirmatory studies (7,8). All these findings have contributed to the ranking of mitotic figure count as a category I prognostic factor by the College of American Pathologists (116), that is, as a factor proven to be of prognostic importance and usefulness in clinical patient management. Mitotic figure count is the oldest measurement of cell proliferation, with a high degree of accuracy, and represents an integral part of histological grade. Furthermore, it is routinely assessable on histological sections/cytological smears, without the need for additional processing or staining procedures. These advantages over the other proliferation indices probably convinced panelists of the most recent NIH-NCI Consensus Development Conferences to support the use of mitotic figure count in clinical practice, alone or in association to the other components of grading systems (116,117). However, if mitotic figure count is used in clinical routine, the standardization of its measurement and the assessment of its intra- and interlaboratory reproducibility is mandatory. In fact, in addition to clinicobiological effectiveness and usefulness, defined as the ability to describe a biological process and its impact on clinical outcome when it influences the choice of therapy, laboratory effectiveness (in terms of the presence of quality assurance programs and methods comparison for any analyte) should also be considered to improve the diagnostic armamentarium in oncology.

Proliferative activity is an example of a biomarker that may be both prognostic and predictive. However, present data are insufficient to draw firm conclusions regarding its predictive role when choosing either endocrine or chemotherapy, but only suggestive of a relationship that should be investigated further in independent adjuvant settings and analyzed with techniques appropriately designed to determine the clinical utility of biomarkers.

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Urokinase-Type Plasminogen Activator and PAI-1

Validated Prognostic Factors for Breast Cancer

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CONTENTS

INTRODUCTION UPA: STRUCTURE, FUNCTION, AND MODE OF ACTION ENDOGENOUS INHIBITORS OF UPA UPA AND PAI-1 IN BREAST CANCER UPA AS A PROGNOSTIC MARKER IN BREAST CANCER PAI-1 AS A PROGNOSTIC MARKER IN BREAST CANCERS TRANSFER OF UPA AND PAI-1 TO THE CLINIC PREDICTIVE VALUE OF UPA AND PAI-1 CONCLUSIONS REFERENCES

SUMMARY

Urokinase plasminogen activator (uPA) is a serine protease causally involved in tumor progression. In vivo, uPA can be inhibited by the serpin inhibitor plasminogen activator inhibitor-1 (PAI-1). However, PAI-1 is a multifunctional protein that can also play a role in cell migration, cell adhesion, angiogenesis, and apoptosis. Multiple singleinstitutional studies have shown that both uPA and PAI-1 are potent

Cancer Drug Discovery and Development: Biomarkers in Breast Cancer: Molecular Diagnostics for Predicting and Monitoring Therapeutic Effect Edited by: G. Gasparini and D. F. Hayes © Humana Press Inc., Totowa, NJ and independent prognostic factors in breast cancer. Recently, this prognostic impact was validated in both a prospective randomized trial and a pooled analysis. As well as being prognostic, high levels of uPA/PAI-1 have been shown to predict for relative resistance to hormone therapy in patients with advanced breast cancer but are associated with an enhanced response to adjuvant chemotherapy in early breast cancer. As uPA and PAI-1 are both prognostic and predictive, assay of these factors has the potential to result in the enhanced management of patients with breast cancer. Measurement of these analytes should thus be now considered for use in the routine management of patients with breast cancer.

Key Words: Urokinase plasminogen activator; PAI-1; cancer; tumor marker; Level 1 evidence.

1. INTRODUCTION

The prognosis of a cancer is dependent primarily on its ability to invade locally and form distant metastasis. The formation of metastasis consists of a series of sequential steps involving processing or remodeling of the extracellular matrix (ECM), local invasion, angiogenesis, intravasation, survival of malignant cells in the circulation, extravasation, and finally growth at a secondary site (for review, *see* ref. 1). To complete the metastatic cascade, malignant cells must also evade apoptosis and resist entering a dormant state. A key molecule causally involved in a number of these steps is the trypsin-like protease, urokinase plasminogen activator (uPA).

2. UPA: STRUCTURE, FUNCTION, AND MODE OF ACTION

The *uPA* gene, which in humans is located on chromosome 10q (10q22) (2), encodes a 53-kDa protein. The protein is initially synthesized as a catalytically inactive single-chain peptide. Conversion to the active form can be brought about, at least in vitro, by a number of proteases such as plasmin, cathepsin B, and cathepsin L (for review, *see* ref. 3). The active form of uPA consists of a two-chain molecule in which the amino terminal A-chain is linked to the B-chain by a single disulfide bond. The A-chain (amino acids 1–158) contains an epidermal growth factor (EGF)-like domain (amino acids 1–49) while the B-chain possesses the catalytic site (3).

As a protease, uPA catalyzes conversion of the zymogen plasminogen to the active plasmin. Unlike uPA, plasmin is a broad-spectrum protease capable of hydrolyzing several different substrates. First, it can promote processing of diverse ECM substrates such as fibrin, fibronectin, and laminin (3). Second, it can activate the precursor forms of specific matrix metalloproteses (MMPs) such as MMP-3, MMP-9, MMP-12, and MMM-13 (4). The formation of the active MMPs allows further processing of the ECM, especially interstitial and type IV collagen. Third, plasmin can activate or release a number of growth factors such as fibroblast growth factor-2 (FGF-2), vascular endothelial growth factor (VEGF), and transforming growth factor- β (TGF- β) (5). These pleiotrophic peptides have the potential to enhance tumour progression by stimulating angiogenesis, cell proliferation and migration.

In vivo, uPA-catalyzed proteolysis occurs while the protease is attached to a membrane-anchored receptor known as uPA receptor (uPAR). uPAR, which is a member of the Ly-6 family of molecules, is a 55–60-kDa glycoprotein (3). It consists of three homologous domains—D1, D2, and D3—and is bound to the cell membrane by a glycosyl phosphatidylinositol (GPI) moiety. The primary binding region in uPA for uPAR resides in the growth factor domain, especially in the sequence containing amino acids 19–32(6). At least four distinct regions in uPAR attach directly to uPA. These sequences included amino acids 13–20 and amino acids 74–84 of domain 1 as well as regions in the putative loop of both domains 2 and 3 (7).

Binding of uPA to its receptor has two main consequences. First, it leads to both enhanced and focused proteolysis. Second, ligand receptor interaction results in signal transduction that involves activation of mitogen-activated protein kinase (MAPK), extracellular regulated kinases (ERK) 1 and 2, and other signaling pathways (8). These signaling pathways allow uPA to stimulate cell migration and mitogenesis, modulate cell adhesion, and prevent cell dormancy (3,8).

3. ENDOGENOUS INHIBITORS OF UPA

uPA activity can be blocked by at least three naturally occurring inhibitors—plasminogen activator inhibitor-1 and -2 (PAI-1, PAI-2) and maspin. These are relatively nonhomologous proteins belonging to the serpin family of protease inhibitors (for review, *see* refs. 9,10). Of the three inhibitors, PAI-1 is thought to be the primary endogenous inhibitor of uPA. The mature form of PAI-1 consists of 381 amino acids and has an apparent molecular mass of 50 kDa. PAI-1 can exist in different conformations (9). The active conformation spontaneously converts into a latent inactive form that in vitro can be reactivated by protein-denaturing agents such as urea and sodium dodecyl sulfate. Alternatively, following inhibition of target proteases, the active conformation is transformed into a center-cleaved inactive form.

As well as binding to uPA, PAI-1 can also associate with a number of different proteins including fibrin, heparin, α_1 acid glycoprotein, and the ECM protein vitronectin. Binding to both vitronectin and α_1 acid glycoprotein stabilizes PAI-1 (10). Furthermore, the interaction with vitronectin allows PAI-1 to modulate cell adhesion by competing with integrins and

uPAR for binding to vitronectin (11). By associating with fibrin, PAI-1 can control dissolution of blood clots (12) while binding to heparin enhances the inhibitory capacity of PAI-1 against thrombin (13).

Recently, PAI-1 was shown to prevent both spontaneous and druginduced apoptosis in a number of different cell lines (14). In contrast to a stable variant of PAI-1, apoptosis was not blocked by a latent form of PAI-1, a stable variant inactivated by specific neutralizing antibodies or a stable variant in complex with uPA. These findings suggest that the inhibitory action of PAI-1 is required for its antiapoptotic effects but that uPAR was not involved (14).

Clearly, PAI-1 can be regarded as a multifunctional protein with several activities in additional to its well-established role in the inhibition of uPA catalytic action. The particular action(s) may depend on factors such as its concentration, its cellular or extracellular localization, and its conformations state.

3. UPA AND PAI-1 IN BREAST CANCER

Multiple studies have shown increased expression of both uPA and PAI-1 in breast carcinomas vis-a vis nonmalignant breast tissue (15-17). The mechanism(s) responsible for this enhanced expression in malignancy are unknown. In breast cell lines in culture, various hormones and growth factors have been shown to modulate uPA expression (Table 1). Other factors known to alter uPA expression in vitro include methylation of gene promoter regions (28) and specific transcription factors, especially members of the ets family (29).

Several groups have related levels of both uPA and PAI-1 in breast cancers to tumor and patient characteristics. In a pooled analysis of 8175 primary breast cancers, uPA levels were not significantly associated with patient age, menopausal status, or lymph node status. Levels however, were positively correlated with tumor grade, negatively with hormone receptors and higher in pT2 and pT4 tumors than pT1 and pT3 samples (*30*).

In contrast to uPA, PAI-1 levels were significantly related to patient age, were higher in postmenopausal than premenopausal women, and higher in lymph node–positive than lymph node–negative cancers. As with uPA, PAI-1 levels were also positively correlated with tumor grade, negatively with hormone receptors and higher in pT2 and pT4 tumors than pT1 and pT3 samples (*30*).

Conflicting reports have been published on the cell type(s) expressing uPA and PAI-1 in breast cancer. Early work using immunohistochemistry suggested that uPA protein was mostly located in malignant cells (15,31,32). Christensen et al. (33), however, reported that uPA immunostaining was present in a number of different cell types, being most intense in macrophages and mast cells but showing moderate intensity in epithelial, fibro-

Factor	Cell line/type	$\uparrow \mathit{or} \downarrow$	Authors (ref.)
Estradiol Estradiol EGF EGF EGF Estradiol FGF-1 FGF-2	Cell line/type ZR-75-1 T47-D MCF-7 SP1 A3a MDA-MB-231 S30 Breast fibroblast Breast fibroblast	$\uparrow or \downarrow$ \uparrow \uparrow \uparrow \uparrow \uparrow \uparrow \uparrow \uparrow	Mangel et al. 1988 (18) Mangel et al. 1988 (18) Pourreau-Schneider et al. 1989 (19) Korczak et al. 1991 (20) Korczak et al. 1991 (20) Long et al. 1996 (21) Levenson et al. (22) Sieuwerts et al. 1999 (23) Sieuwerts et al. 1999 (23)
IGF-1 IGF1 Progesterone Amphiregulin Heregulin	Breast fibroblast MDA-MB-231 MDA-MB-231 ^a MCF-7 LM3	$\stackrel{\downarrow}{\uparrow} \stackrel{\downarrow}{\rightarrow} \stackrel{\downarrow}{\rightarrow}$	Sieuwerts et al. 1999 (23) Dunn et al, 2000 (24) Lin et al. 2001 (25) Silvy et al. 2001 (26) Puricelli et al. 2002 (27)

Table 1 Factors Shown to Regulate Expression of uPA Levels in Cell Lines Derived from Breast Cancers

 \uparrow , Increase expression; \downarrow , decreased expression.

^aTransfected with progesterone receptor.

blasts, and endothelial cells. Kennedy et al. (34) also found uPA staining in both stromal and epithelial cells, although the predominant location was in stromal cells. Finally, Nielsen et al. (35) identified uPA immunoreactivity in myofibroblasts and macrophages in all 25 breast cancers examined. In 12 cases, endothelial cell staining was also seen. Only three cases exhibited cancer cell positivity and in each, only a small subpopulation of cells stained. These findings of Nielsen et al. (35) are consistent with previous *in situ* hybridization studies from the same group showing that mRNA for uPA was located almost exclusively in fibroblasts (36). The finding of uPA in stromal cells suggests that these cells synergize with cancer cells to promote metastasis.

Disagreement also exists on the location of PAI-1 in breast cancer. For example, Reilly et al. (16) detected PAI-1 protein in both endothelial and cancer cells. Similarly, Christensen et al. (33) found PAI-1 in endothelial and cancer cells, but also located it in stromal cells. In contrast, Jankum et al. (32) reported that PAI-1 was present mostly in malignant cells.

Possible reasons for these conflicting results on the location of uPA and PAI-1 in breast carcinomas include use of different methods of processing tissue (fresh vs formalin-fixed and paraffin-embedded tissue), different pretreatments of tissue, use of antibodies of different specificities, and possible failure in some studies to rigorously exclude nonspecific staining.

Table 2 Evidence Implicating uPA in Invasion and Metastasis

Positive correlations are found between levels of uPA in both cell lines and animal tumors and metastatic potential.

Inhibition of uPA activity (e.g., by inhibitors or antibodies) or uPA expression (e.g., by antisense oligonucleotides) suppresses metastasis in model systems.

Transfection of cell lines with cDNA for uPA enhances the metastatic phenotype of recipient cells.

Prevention of uPA from binding to uPAR decreases metastasis in model systems. Tumors in uPA deficient mice undergo less progression than in control wildtype mice.

Data summarized from refs. 37,38.

4. UPA AS A PROGNOSTIC MARKER IN BREAST CANCER

Studies with multiple model systems have shown that uPA is causally involved in metastasis (37,38) (Table 2). Originally, uPA was thought to play a role in these processes simplify by degrading the ECM, thus permitting local invasion and ultimately the formation of distant metastasis. The degradation of the ECM is now thought to be highly specific with limited substrate cleavage, that is, protein processing. However, in addition to protein processing, it is now clear that uPA has other actions allowing it mediate cancer cell dissemination (3). These activities include its ability to enhance angiogenesis, stimulate both cell proliferation and migration, and inhibit dormancy (for review, *see* refs. 3,8).

Because the formation of distant metastasis is the principal cause of mortality in patients with cancer and uPA is a critical mediator of the process, the latter should be a good candidate for investigation as a prognostic marker (39). Furthermore, the positive correlations found between uPA levels and metastatic ability in both cell lines and animal cancers (37) might also be expected to apply to human cancers.

Duffy et al. (40,41) first reported that breast cancer patients with high levels of uPA activity in their primary cancer exhibited both a shorter disease-free interval and shorter overall survival than those with low activity levels. These preliminary results have now been confirmed by approx 20 independent groups, worldwide (Tables 3 and 4). These singlecenter studies showed that the prognostic value of uPA was:

- Independent of that provided by the traditional prognostic factors for this disease such as tumor size, tumor grade, or axillary node status.
- Stronger than that of other biological factors such as estrogen receptor, progesterone receptor, *HER2*, EGFR, p53, and Nottingham Index (60,62,63).

Authors (ref.)	No.	R or P study	Comments
Duffy et al. 1988 (40)	52	R	
Janicke et al. 1989 (42)	104	Р	
Spyratos et al. 1992 (43)	319	R	
Cook et al. 1992 (44)	48	NS	
Foekens et al. 1992 (45)	671	R	
Grondahl-Hansen et al. 1993 (46)	119	NS	Only premenopausal patients
Bouchet et al. 1994 (47)	316	NS	
Sumiyoshi et al. 1995 (48)	80	NS	
Ferno et al. 1997 (49)	688	NS	
Shiba et al. 1997 (50)	226	Р	
Umeda et al. 1997 (51)	73	R	
Koop et al. 1998 (52)	429	R	
Peyrat et al. 1998 (53)	634	NS	
Kim et al. 1998 (54)	130	Р	
Tetu et al. 1998 (55)	575	R	
Eppenberger et al. 1998 (56)	305	NS	Only node-negative patients
Kute et al. 1998 (57)	116	NS	Only node-negative patients
Broet et al. 1999 (58)	1245	R	-
Konecny et al. 2001 (59)	587	R	
Malmstrom et al. 2001 (60)	237	Р	Only premenopausal node-negative patients
Meo et al. 2002 (61)	196	NS	Only node-negative patients

 Table 3

 Different Groups Showing a Prognostic Value for uPA in Breast Cancer

Many of the above-listed groups have published follow-up studies with greater numbers of patients and longer follow-up. Only the original study is cited above. P, Prospective study; R, Retrospective study; NS, not stated.

- Independent of the cutoff point used for separating patients with low and high levels of uPA, that is, whether the median, tertile, or quartile value was used (64). uPA was also predictive when used as continuous variable (64).
- Prognostic in axillary node-negative patients (Table 4), including node-negative patients who did not receive systemic adjuvant therapy (65).

in Axillary Node-Negativ	ve Breast Cancer	r Patients
Authors (ref.)	No.	R or P study
Janicke et al. 1990 (15)	50	Р
Foekens et al. 1992 (45)	272	R
Duffy et al. 1994 (66)	75	R
Ferno et al. 1997 (49)	265	NS
Peyrat et al. 1998 (53)	634	NS
Kim et al. 1998 (54)	130	Р
Eppenberger et al. 1998 (56)	305	NS
Malmstrom et al. 2001 (60)	237	Р
Meo et al. 2002 (61)	196	NS

Table 4 Different Groups Showing a Prognostic Value for uPA in Axillary Node-Negative Breast Cancer Patients

Some of the above-listed groups have published follow-up studies with greater numbers of patients and longer follow-up. Only the original study is cited above. P, Prospective study; R, retrospective study; NS, not stated.

5. PAI-1 AS A PROGNOSTIC MARKER IN BREAST CANCERS

Intuitively, it might be expected that high levels of an inhibitor of uPA in cancer tissue would correlate with a low probability of metastasis and thus with good outcome. Paradoxically, however, high levels of PAI-1 in breast cancer are also strongly and independently associated with poor outcome (Tables 5 and 6). Furthermore, the prognostic information available from PAI-1 is additional to that of uPA. Thus, the combined measurement of both proteins results in enhanced prognostic data over that available from either factor alone (65).

Why high levels of PAI-1 correlate with adverse prognosis is not clear. Possible explanations include:

- A critical concentration of PAI-1 is necessary to prevent excessive degradation of the ECM by uPA during cancer invasion. Excessive breakdown of the matrix could leave insufficient substrate for migration of cancer cells.
- PAI-1 is necessary for angiogenesis (72,73), which in turn is essential for malignant cells to gain access to the circulation and for progression at both primary and metastatic sites.
- PAI-1 can modulate both cell adhesion and migration (11) and as result may accelerate the metastatic process.
- PAI-1 can inhibit apoptosis (14). For malignant cells to complete the metastatic process, they must survive and evade apoptosis. Resistance

Authors (ref.)	No.	R or P study	Comments
Janicke et al. 1991 (67)	102	Р	
Grondahl-Hansen et al. 1993 (46)	72	NS	Only postmenopausal patients
Bouchet et al. 1994 (47)	314	NS	•
Foekens et al. 1994 (68)	657		
Sumiyoshi et al. 1995 (48)	80	NS	
Duggan et al. 1997 (69)	148	R	
Eppenberger et al. 1998 (56)	305	NS	Only node-negative patients
Kute et al. 1998 (57)	135	NS	Only node-negative patients
Knoop et al. 1988 (52)	429	R	L
Kim et al. 1988 (54)	130	Р	Only node-negative patients
Billgren et al. 2000 (70)	1851	NS	•
Konecny et al. 2001 (59)	587	R	
Meo et al. 2002 (61)	196	NS	Only node-negative patients

 Table 5

 Different Groups Showing a Prognostic Value for PAI-1 in Breast Cancer

Many of the above-listed groups have published follow-up studies with greater numbers of patients and longer follow-up. Only the original study is cited above. P, Prospective study; R, retrospective study; NS, not stated.

Table 6
Different Groups Showing a Prognostic Value for PAI-1
in Axillary Node-Negative Breast Cancer Patients

Authors (ref.)	No.	R or P study
Janicke et al. 1993 (71)	101	Р
Foekens et al. 1994 (68)	272	R
Bouchet et al. 1994 (47)	146	NS
Kim et al. 1998 (54)	130	Р
Meo et al. 2002 (61)	96	NS

Many of the above-listed groups have published follow-up studies with greater numbers of patients and longer follow-up. Only the original study is cited above. P, Prospective study; R, retrospective study; NS, not stated. to apoptosis might therefore be expected to increase the likelihood of forming a distant metastasis. Evasion of apoptosis could also confer resistance to specific therapies (74), which in turn could result in a more adverse outcome.

6. TRANSFER OF UPA AND PAI-1 TO THE CLINIC

Prior to entering routine clinical use, new markers should be rigorously evaluated with respect to both analytical and clinical performance. Several research and commercially available enzyme-linked immunosorbent assay (ELISA) kits have now been described for the measurement of uPA and PAI-1 concentration (75). A number of these have been subjected to detailed evaluation with the following conclusions (75,76).

- All the assays tested that were designed for the measurement of uPA in tissue extracts had adequate sensitivity for application in breast cancer extracts.
- Within assay precision for all the assays investigated was satisfactory.
- All assays investigated displayed an acceptable degree of parallelism following dilution of tissue extracts.
- Although the absolute level of uPA measured varied with the different methods, in general, good correlations were found between the different assays.
- Higher yields of uPA were obtained with a Triton X-100–containing buffer than with a detergent-free buffers.
- Some uPA and PAI-1 kits (American Diagnostica, Stamford, CT) were subsequently investigated in external quality assurance studies and shown to perform in a satisfactory manner (76). For example, in a German multicenter clinical trial in which six different laboratories all used these kits, the interlaboratory coefficient of variation (CV) varied between 6.2 and 8.2 for uPA and between 13.2 and 16.6 for PAI-1. Expressing results as nanograms per milligram of protein however, led to a substantial increase in the interlaboratory variation, that is, CVs of 10.8–20.4 for uPA and 15.5–23.6 for PAI-1.

Clinical validation of a new marker, requires validation in either a large randomized prospective trial in which evaluation of the marker is the primary objective of the study or a meta-analysis/pooled analysis of small-scale prospective or retrospective trials (77). Recently, the prognostic impact of uPA/PAI-1 in breast cancer patients was confirmed using both these types of Level 1 Evidence studies (30,78).

The prospective randomized investigation was a multicenter study containing almost 600 patients carried out in Germany (78). In this trial, nodenegative breast cancer patients with low levels of uPA and PAI-1 were monitored but received no systemic adjuvant chemotherapy. On the other hand, patients with high levels of the protease and/or its inhibitor were randomized to receive adjuvant CMF treatment or to be observed. Following an interim analysis after 32 mo of follow-up, patients with low levels of both proteins had an estimated 3-yr recurrence rate of 6.7% whereas those with high concentrations of the protease and/or its inhibitor had a recurrence rate of 14.7% (p = 0.006).

Multivariate analysis showed that the prognostic impact of uPA/PAI-1 was independent of tumor grade, tumor size, surgical treatment of the primary cancer, and steroid receptor status. Although tumor grade was a stronger predictor of outcome than uPA/PAI-1, if grade was used to classify risk, only 10% of the node-negative patients would be regarded as having a low risk of recurrence, that is, those with grade 1 disease. In contrast, based on uPA/PAI-1, 56% would be considered at low risk of developing disease recurrence (78).

The second type of Level 1 Evidence study to have validated the prognostic significance of uPA and PAI-1 in breast cancer involved a pooled analysis of 18 different data sets containing a total of 8377 patients (*30*). Of these 18 studies, 11 were previously published while seven contained unpublished data. Raw data from all the 18 studies were used in the statistical analysis. Following a median follow-up of 79 mo, both uPA and PAI-1 were found to independent prognostic factors. Although less potent than axillary nodal status, both uPA and PAI-1 were stronger predictors of outcome than tumor size, tumor grade, hormone receptor status, or patient age. In the node-negative patients, uPA and PAI-1 were the strongest predictors of both disease-free interval and overall survival. Significantly, both uPA and PAI-1 were also prognostic in the subgroup of node-negative patients who did not receive systemic adjuvant therapy (*30*).

To the author's knowledge, uPA and PAI-1 are the first biological factors for which prognostic value was validated using either a prospective randomized trial or a pooled analysis. The results of the two studies clearly show that node-negative breast cancer patients have a low risk of disease relapse and consequently could avoid the side effects and financial costs of adjuvant chemotherapy. Assay of uPA and PAI-1 should therefore be now considered for routinely determining prognosis in axillary node-negative breast cancer patients.

7. PREDICTIVE VALUE OF UPA AND PAI-1

According to Clark (79), prognostic factors are necessary not only for predicting patient outcome but also to indicate which patients are likely to benefit from specific therapies. In a pilot study containing 235 patients with recurrent breast cancer, Foekens et al. (80) reported that women with uPA-negative tumors exhibited a better response to tamoxifen therapy than those

with uPA-positive tumors. This predictive impact of uPA appeared to be independent of steroid receptor status. When patients were stratified by receptor concentration, however, the relationship between high levels of uPA and resistance to tamoxifen was observed only in the subgroup with intermediate levels of estrogen receptors (ERs) and progesterone receptors (PgRs) (i.e., >10 fmol/mg of protein for both receptors with at least 1 being not more than 75 fmol/mg). High levels of PAI-1 were also associated with reduced benefit from tamoxifen but the predictive effect of the inhibitor was less pronounced than that of the protease (*80*).

While high levels of uPA/PAI-1 have been reported to predict for resistance to hormone therapy in patients with advanced breast cancer, recent data suggest that elevated levels of these proteins correlate with an enhanced response to adjuvant chemotherapy. In the German prospective randomized trial referred to in the preceding (78), administration of chemotherapy to patients with high levels of uPA and/or PAI-1 reduced the relative risk of recurrence by approx 44% (relative risk = 0.56, *p*, NS). However, if patients who violated the study protocol were excluded from the analysis, the benefit of CMF became more pronounced (p = 0.016, relative risk = 0.27). This finding suggested that lymph node–negative breast cancer patients with high levels of uPA/PAI-1 benefit from adjuvant CMF.

Consistent with these findings, Harbeck et al. (65), in a single-institution prospective study (n = 761), found that while uPA/PAI-1 predicted outcome in patients who did not receive systematic adjuvant therapy, the prognostic impact was lost in patients treated with either adjuvant chemotherapy or adjuvant hormone therapy. Again, this study suggested a benefit from adjuvant treatment in patients with high uPA/PAI-1 levels. More direct evidence of a predictive role for uPA/PAI-1 has come from a large two-center study (n = 3424), which showed that breast cancer patients with high levels of these proteins benefited more strongly from adjuvant chemotherapy than those with low levels (81).

These preliminary findings suggest that in advanced breast cancer, high levels of uPA and PAI-1 predict for resistance to tamoxifen therapy. On the other hand, in early breast cancer, patient with high levels appear to derive an enhanced benefit from CMF-based chemotherapy. These promising results should now be confirmed in high-powered prospective randomized trials.

8. CONCLUSIONS

uPA and PAI-1 are among the few tumor markers whose clinical value has been validated using Level 1 Evidence data, and to the author's knowledge are the only markers to have been validated using two different types of such studies. Because assays for these markers have also been shown to perform satisfactorily in EQA trials, both should now be ready for routine clinical use. In the clinic, the immediate application of these markers is likely to be in selecting node-negative breast cancer patients who do not need or are unlikely to benefit from adjuvant chemotherapy, that is, patients with low levels of uPA and/or PAI-1.

In addition to being prognostic, uPA and PAI-1 have been reported to predict for likely response or resistance to therapy in patients with breast cancer. As mentioned earlier, emerging data suggests that high levels of these proteins are associated with resistance to hormone therapy in advanced breast cancer but correlate with enhanced benefit from adjuvant CMF therapy in early breast cancer.

As well as being able to predict for likely response to specific therapies, uPA can also be a direct target for new forms of anticancer therapies. Data from model systems show that either inhibition of uPA activity or blocking uPA from binding to its receptor reduces tumor growth and metastasis (82). Recently, a serine protease inhibitor known as WX-UK1 (Wilex AG, Munich, Germany) and which is directed against uPA and plasmin, entered clinical trials. It might be expected that the tumors most likely to respond to uPA-directed therapy will be those expressing high levels of uPA. Thus, like ER and HER-2, uPA/PAI-1 has the potential to be both prognostic and predictive as well as a target for therapy.

In conclusion, clearly a knowledge of uPA and PAI-1 levels have the potential to result in the enhanced management of patients with breast cancer. Assay of these factors is therefore likely to be introduced into routine clinical practice in the near future. A limiting factor of the traditional assays for uPA and PAI-1 was the requirement for at least 100 mg of tumor tissue. Recently, however, a microassay for the detection of uPA/PAI-1 has been described (83). This assay utilizes one or two breast tissue core biopsies or 5-10 of $90-\mu$ M-thick cryosections of a frozen tumor block. The availability of this new assay should allow the determination of uPA and PAI-1 in small breast cancers.

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8

Predictive Value of c-*erb*-B2 for Endocrine Therapy and Chemotherapy in Breast Cancer

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CONTENTS

INTRODUCTION C-*erb*-B2 as a Predictive Factor FOR RESPONSE TO ENDOCRINE MANIPULATIONS C-*erb*-B2 as a Predictive Factor FOR RESPONSE TO CHEMOTHERAPY C-*erb*-B2 as a Predictive Factor FOR RESPONSE TO TRASTUZUMAB CONCLUSIONS REFERENCES

SUMMARY

c-*erb*-B2, also designated *HER-2/neu* or c-*neu*, is a tyrosine kinase proto-oncogene that may be overexpressed or amplified in 20–40% of breast tumors. Data reported to date suggest that the proto-oncogene may be a predictive factor of response to several therapies that are commonly used to treat breast cancer. Tumors that are both hormone receptors and c-*erb*-B2 positive may be associated with a relative resistance to tamoxifen. Preliminary data suggest that aromatase inhibitors may be more effective in such tumors. Available data suggest that women with c-*erb*-B2–positive tumors may have a

Cancer Drug Discovery and Development: Biomarkers in Breast Cancer: Molecular Diagnostics for Predicting and Monitoring Therapeutic Effect Edited by: G. Gasparini and D. F. Hayes © Humana Press Inc., Totowa, NJ preferential response to anthracycline-containing regimens compared to cyclophosphamide, methotrexate, 5-fluorouracil (CMF)-like regimens. Anthracycline-based therapy should be recommended to women with c-*erb*-B2–positive breast cancer unless there is a contraindication to the administration of this group of agents. However, other regimens should not be withheld from women who cannot receive an anthracyline. Current clinical data also suggest a possible sensitivity to taxane-based therapy, especially when combined with an anthracycline. Trastuzumab-based therapy should be considered as first line therapy for women with metastatic breast cancer with c-*erb*-B2–positive disease who are hormone receptor negative, or those with hormone receptor–positive disease who have progressed on endocrine treatments. Adjuvant trastuzumab should be considered in women with high-risk primary breast cancer.

Key Words: c-*erb*-B2; *HER-2/neu*; predictive factor; hormone therapy; chemotherapy; trastuzumab.

1. INTRODUCTION

c-erb-B2, also designated HER-2/neu or c-neu, is a tyrosine kinase protooncogene that belongs to the epidermal growth factor receptor (EGFR) family. The EGFR family includes four members designated HER-1 (EGFR), HER-2, HER-3, and HER-4. The receptor consists of a transmembrane domain connected to an extracellular domain and an intracellular domain. When a ligand binds to the extracellular domain, the receptor homoor heterodimerizes with another HER receptor and the intracellular tyrosine kinase domain is activated via phosphorylation. A cascade of proteinto-protein interactions is initiated, and a signal of growth and proliferation is transmitted to the nucleus. Of note, a ligand to c-erb-B2 has not been identified. Up to 20-40% of breast cancers overexpress or amplify c-erb-B2. Despite two decades of investigation and hundreds of publications, the role of c-*erb*-B2 as a prognostic and predictive factor in breast cancer is still in question. Indeed, members of the American Society of Clinical Oncology (ASCO) Tumor Marker Expert Panel, the NIH Consensus Panel, and the St. Gallen International Consensus Panel determined that c-erb-B2 status should not be used to alter treatment recommendations for individual patients (1-3).

Prior to discussing individual studies that evaluated the predictive role of c-*erb*-B2, it is important to review some of the reasons for the controversial role of the marker. Difficulties assessing the true predictive role of a marker such as c-*erb*-B2 stem from heterogeneity in study design and patient population as well as variability in methods of marker evaluation (Table 1). Generally, anecdotal or preclinical data that suggest a marker may have a

Table 1

Reasons for the Controversial Role of c-*erb*-B2 as a Predictive Marker for Treatments of Breast Cancer

Heterogeneity in study design, patient population, and follow-up
Retrospective analyses of archival tissue
No distinction between prognostic and predictive role
Include patients with many Tumor–Node–Metastasis (TNM) stages
Include patients with many treatments
Treatments may be suboptimal by today's standards
Not all trial patients included in marker study
Patients' outcomes not known
Heterogeneity of methods of marker detection
Assays used detect different abnormal processes
Gene amplification
RNA amplification
Protein overexpression
Evaluated tissue or circulating marker levels
Different specimen preparation
Different reagents
Different cutoff level

prognostic or predictive role in a specific cancer led to retrospective analyses of samples that may have been collected through previous clinical trials or simply from tissue banks. The patient population in a marker study may be heterogeneic and information regarding therapy, follow-up, specimen selection, or statistical analysis is not usually prospectively determined. Such studies are hypothesis-generating and designated level of evidence (LOE) III or IV studies (4). Results of LOE III or IV studies must be further tested and validated before a marker is ready for clinical use. After establishing a possible role, LOE I evidence (i.e., evidence from a single highpowered prospective study that is specifically designed to test the marker or evidence from meta-analysis and/or overview of lower evidence studies) is required prior to accepting the marker for clinical use (4). Unfortunately, most investigations that evaluated the role of c-erb-B2 in predicting response to breast cancer treatments include LOE III or IV studies. Several studies have evaluated c-erb-B2 in samples that have been collected through a prospective trial that was designed to test therapeutic hypothesis with a marker study as a secondary objective defined in the protocol (LOE II studies). Unfortunately, despite prospective collection of samples for the marker study, samples are often available only from a subgroup of patients who were included in the clinical trial.

Heterogeneity in study design also contributes to the confusion of whether a marker is prognostic, predictive, or both. A true prognostic marker reflects tumor biology and, in the absence of treatment, its presence may suggest improved long-term outcomes such as disease free and/or overall survival (positive prognostic marker) or worse outcomes (negative prognostic marker). To evaluate the true prognostic role of a marker, absence of systemic treatment is desired. In contrast, the presence or absence of a predictive marker is associated with sensitivity or resistance to specific therapy. To evaluate the predictive role of a marker, outcomes such as response rates, disease-free survival, and overall survival may be calculated for study participants with marker-positive vs marker-negative disease. The presence of the estrogen receptor (ER) is an example of a strong predictive marker for response to endocrine manipulations that is also a weak positive prognostic marker. Current data suggest that c-*erb*-B2 may be a weak negative prognostic factor and will not be reviewed in this chapter (5).

It is also difficult to compare results of individual marker studies owing to the heterogeneity in methods used to analyze the marker. While most early studies of c-*erb*-B2 used immunohistochemistry (IHC) to detect protein expression, others evaluated amplification of the oncogene. Even when the same method is used, different antibodies or cutoff levels may be utilized (Table 1). In this chapter, studies that assessed the predictive role of c-*erb*-B2 to endocrine therapies and to chemotherapies are discussed with an emphasis on studies with high LOE. In this review, c-*erb*-B2–positive tumors are referred to tumors that overexpress or amplify the oncogene using the cutoff levels and criteria defined in the individual study.

2. c-*erb*-B2 AS A PREDICTIVE FACTOR FOR RESPONSE TO ENDOCRINE MANIPULATIONS

Tumors from more than 50% of women diagnosed with breast cancer will express the ER and/or progesterone receptor (PgR). Most of hormone receptor-positive women will be offered endocrine manipulations at some point during the treatment course. The most common endocrine therapy administered to women with breast cancer is tamoxifen. Aromatase inhibitors are commonly administered as first- or second-line therapy for menopausal women with metastatic breast cancer. Based on the results from the Arimidex, Tamoxifen, Alone or in Combination (ATAC) Trial, the United States Food and Drug Administration recently approved the aromatase inhibitor anastrozole for adjuvant therapy for women with hormone receptor-positive breast cancer (6). Other studies support the use of the aromatase inhibitor letrozole following 5 yr of tamoxifen, or exemestane for 2-3 yr following 2-3 yr of tamoxifen for a total of 5 yr (7,8). Based on these recent reports, aromatase inhibitors are likely to have an increasing role in the adjuvant treatment of hormone receptors positive postmenopausal women (9). Ovarian ablation may be employed in premenopausal or perimenopausal women with hormone receptor–positive disease (10). Initial reports from preclinical studies and retrospective clinical trials suggested that c-*erb*-B2– positive tumors may be associated with relative resistance to hormone therapies. However, the data are inconclusive. Reports from recent studies may also suggest that the presence of c-*erb*-B2 may not confer resistance to all endocrine therapies.

2.1. Preclinical Data

Breast cancer cell lines that overexpress c-erb-B2 or those that have been transfected with the oncogene are tamoxifen resistant (11-13). Tamoxifen inhibits the growth of parental MCF-7 xenografts but not the growth of c-erb-B2-transfected MCF-7 xenografts (11-13). One hypothesis that may explain tamoxifen resistance is that while tamoxifen inhibits ER-stimulated growth, the cells may be activated through the c-erb-B2 signal transduction pathway. Other data suggest that tamoxifen, unlike estrogen, may up-regulate the expression of c-erb-B2. Indeed, MCF-7 cells transfected with c-erb-B2 are associated with mitogen-activated protein (MAP) kinase hyperactivity and tamoxifen resistance. When c-erb-B2 and MAP kinase are blocked, tamoxifen sensitivity is restored (14). It is also possible that hormone receptor-positive tumors that also overexpress c-erb-B2 have lower expression of the hormone receptors compared to similar tumors that are c-erb-B2 negative (15). Thus, the c-erb-B2-positive tumors may be less responsive to the hormone therapies simply because of a lower hormone receptor content.

These preclinical studies suggest that there may be a "cross-talk" between the signal transduction associated with c-*erb*-B2 and that of the ER. Tumors that overexpress or amplify c-*erb*-B2 may be hormone independent and may indeed be relatively resistant to tamoxifen. However, these preclinical studies do not provide data regarding the predictive role of c-*erb*-B2 to hormonal manipulations that deprive the cells of estrogen without a direct association with the ER, such as aromatase inhibitors or ovarian suppression.

2.2. Studies of Hormone Therapies in Metastatic Breast Cancer

Several studies evaluated the predictive role of *c-erb*-B2 to hormonal therapies in metastatic breast cancer. Because women with metastatic breast cancer require effective therapy, randomized clinical trials with a control group are not available. However, one can assume that in the absence of treatment the response rate of the tumor will be nil. Then, with the use of any therapy, response rates and other outcomes can be compared between women with tumors that are *c-erb*-B2 positive vs those whose tumors are *c-erb*-B2 negative.

Most studies that investigated tissue c-erb-B2 status using IHC in patients with metastatic breast cancer reported that overexpression of the marker was associated with relative resistance to tamoxifen. In addition to differences in patient population and methods of c-erb-B2 detection, the studies are mostly small and may have included women with hormone receptorpositive and -negative tumors who may have received one of several hormone therapies. In one large study, women received first line tamoxifen (n = 211) or ovarian ablation (n = 30). Response rate and time to progression (TTP) were statistically significantly worse in women who overexpressed c-erb-B2 compared to women with c-erb-B2-negative disease (16). When the analysis was restricted to the 189 women whose tumors were ER positive, TTP (5.5 vs 11.2 months, p < 0.001) and response rate (24% vs 64%, p = 0.05) were worse in c-*erb*-B2–positive patients. In a study of 104 women who received tamoxifen and 22 who received high-dose progestin or other hormone therapies, response rates were worse in c-erb-B2-positive women as was disease-free survival (DFS) (17). A smaller study included samples from 65 women treated with tamoxifen or aminoglutetamide and hydrocortisone. Response rates were 7% for c-erb-B2-positive patients compared to 37% for c-erb-B2-negative patients (18).

Studies that were restricted to ER-positive women who received tamoxifen only revealed no association between c-*erb*-B2 status and response to tamoxifen. In South West Oncology Group (SWOG) study 8228, evaluation of 205 samples of 349 study participants revealed no significant difference in response rate (54% and 57%), DFS (6 and 8 mo), or OS (29 and 31 mo) between women with c-*erb*-B2–positive and –negative disease (19). In a smaller study that evaluated circulating extracellular domain (ECD) of c-*erb*-B2, significant difference in response to tamoxifen was not observed regardless of c-*erb*-B2 status (20). Although both women with ER-positive or -negative disease were included in the initial analysis, the results remained insignificant even when the analysis was restricted to ER-positive patients.

Early reports suggested that c-*erb*-B2 positivity may also predict resistance to endocrine treatments other then tamoxifen. Response rates to treatments with megestrol acetate, fadrazole, and droloxifene were worse in women with elevated levels of ECD-c-*erb*-B2 compared to those who did not have elevated levels of the circulating marker (21,22). These results supported the hypothesis that c-*erb*-B2 may confer a worse response to hormone therapies. However, while ECD shedding closely correlates with tumors that overexpress or amplify c-*erb*-B2 (high specificity), not all women whose tumors overexpress c-*erb*-B2 will have detectable circulating levels of the ECD of the marker (23). At the same time, ECD–c-*erb*-B2 concentrations can be more precisely quantified compared to IHC, and the degree of marker elevation may be predictive of response. Large studies correlating c-*erb*-B2 status and response to the aromatase inhibitor letrozole have demonstrated conflicting results. Colomer evaluated ECD–c-*erb*-B2 levels in 223 women with hormone receptor–positive metastatic breast cancer who received second-line hormone therapy with letrozole. Time to failure (TTF) was shorter for women with elevated concentrations of ECD–c-*erb*-B2 compared with those whose levels were below the cutoff value (24). Similar results were observed in a study of 711 women who received megestol acetate, fadrazole, or letrozole. Women with elevated concentrations of ECD–c-*erb*-B2 had a worse response rate (23% vs 45%, p < 0.0001), TTP (3 vs 6 mo, p < 0.0001), and overall survival (17.2 vs 29.6 mo, p < 0.0001) compared to those whose concentrations were below the cutoff (25).

Outcomes of women with c-erb-B2-positive or -negative breast cancer were evaluated in two randomized clinical trials of tamoxifen vs letrozole. In the first study, postmenopausal women with hormone receptor-positive locally advanced breast cancer received neoadjuvant therapy with letrozole or tamoxifen. Overall, response rate was greater for patients who received letrozole compared to those who received tamoxifen (60% and 41%, respectively, p = 0.004). In a post hoc analysis, the presence of c-*erb*-B2 (IHC) predicted for improved response to letrozole compared to tamoxifen (Table 2). Response rate to letrozole was 69% and 53% for women with c-erb-B2positive and -negative disease (odds ratio [OR] for response 1.93, 95% CI 0.63-5.88, p = 0.25), compared to much lower response rates to tamoxifen of 17% and 40% for women with marker-positive and -negative disease, respectively (OR 0.31, 95% CI 0.10–0.97, p = 0.045). The odds ratio for response to letrozole vs tamoxifen for women whose tumors overexpressed *erb*-B1 or *erb*-B2 was 28 (response rate 88% vs 21%, respectively, p = 0.0004) compared to an odds ratio of 1.7 (response rate 54% vs 42%, p = 0.078) in c-erb-B1- or -2-negative patients (26). Similar results were reported in a randomized trial comparing neoadjuvant anastrozole vs tamoxifen. Women with both ER and c-erb-B2 positive tumors were much more likely to respond to anastrozole compared to tamoxifen (27). While intriguing, these data are a result of an unplanned analysis of a small number of patients whose tumors were c-erb-B2 positive and validation is required in larger studies. In addition, it is not known whether the greater response rates in the c-erb-B2-positive women will translate to improved DFS and/or OS. In another large study of tamoxifen vs letrozole as first line therapy for metastatic breast cancer, letrozole-treated women had improved outcomes compared to tamoxifen-treated women regardless of ECD-c-erb-B2 concentrations. Importantly, women without elevated ECD-c-erb-B2 who were treated with tamoxifen had substantially better outcomes compared to women with elevated ECD-c-erb-B2 who were treated with either tamoxifen or letrozole (Table 2). Thus, even if women with c-erb-B2-

		a. At	a. Advanced Breast Cancer	cer		
	Patient population and number in marker study	Treatment arms (randomized, unless otherwise	Respon arm I vs arn	Response rate arm 1 vs arm 2 (p value)	Time to p arm I vs arr	Time to progression arm 1 vs arm 2 (p value)
Author	(% c - erb - B2 - positive)	-	c-erb-B2-positive	c-erb-B2-positive c-erb-B2-negative	c-erb-B2-positive	c-erb-B2-positive c-erb-B2-negative
Ellis (26)	LABC, first line, ER or PR+ 250 (14%)	1. Tamoxifen 20 mg 2. Letrozole 2.5 mg	17% vs 69%	40% vs 53%	NR	NR
Lipton (84)	LABC, metastatic, first line 562 (29%)	1. Tamoxifen 20 mg 2. Letrozole 2.5 mg	13% vs 17% (0.4507)	26% vs 39% (0.078)	3.3 vs 6.1 mo ^a (0.0596)	8.5 vs 12.2 mo ^a (0.0019)

Table 2 Randomized Clinical Trials of Endocrine Treatments

	Patient population and number in marker study		HR disease- (95% CI, arm I v	HR disease-free survival (95% CI, p value) arm I vs arm 2	HR overa (95% CI arm 1 v	HR overall survival (95% CI, p value) arm 1 vs arm 2
Author	(% c-erb-B2-positive)	Treatment arms	c-erb-B2-positive	c-erb-B2-negative	c-erb-B2-positive	c-erb-B2-positive c-erb-B2-negative
Carlamango/ (GUN 1) (28)	N-, ER+/-, no CT 145 (30%)	 Tamoxifen × 2 yr No therapy 	51% vs 63% ^b (0.3)	82% vs 54% ^b (0.003)	57% vs 82% ^b (0.03)	$86\% vs 68\%^b$ (0.09)
Muss, Berry (CALGB 8541) (30,58)	N+, 3 dose level CAF 650 (24%)	 Tamoxifen × 5 yr No tamoxifen (not randomized) 	0.68 (0.42–1.1, 0.12)	0.61 (0.47–0.79, 0.0001)	$\begin{array}{ccc} 0.61 & 0.70 \\ (0.47-0.79, 0.0001) & (0.41-1.18, 0.18) \end{array}$	0.64 (0.47–0.87, 0.0037)
Stal (34)	ER+ 405 (11%)	1. Tamoxifen \times 5 yr 2. Tamoxifen \times 2 yr	2.0 (0.49–7.9)	0.58 ($0.37-0.90$)	NR	NR
Love (85)	N+/-, ER+, no CT 282 (26%)	 Ovarian ablation and tamoxifen No therapy 	0.37 (0.26-0.89, 0.047)	0.48 (0.31–0.71, 0.019)	0.26 (0.07–0.92, 0.038)	0.68 (0.32-1.42, 0.30)

b. Early Breast Cancer (Adjuvant)

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positive disease may be more responsive to aromatase inhibitors compared to tamoxifen, their outcomes following letrozole treatment may still be worse than outcomes of women whose tumors are c-*erb*-B2 negative, reflecting the prognostic role of the marker.

Taken together, the results of studies in advanced breast cancer suggest that women whose tumors were c-*erb*-B2 positive may or may not be relatively resistant to tamoxifen. Whether c-*erb*-B2–positive women may derive greater benefit from aromatase inhibitors compared to tamoxifen is also not clear.

2.3. Studies of Hormone Manipulations in the Adjuvant Setting

Samples for c-erb-B2 analysis were available from only a handful of studies that compared adjuvant tamoxifen or other endocrine therapies to no treatment. The Gruppo Universitario Napoletano (GUN-1) was the first large randomized clinical trial that suggested that c-erb-B2 may be a negative predictive factor for response to tamoxifen. From 1978 to 1983, 433 women whose ER status was unknown were randomly assigned to 30 mg of tamoxifen daily for 2 yr or to no hormone therapy. Of those, 173 women were node negative and did not receive adjuvant chemotherapy, and tumors from 145 (84%) were available for c-erb-B2 analysis by IHC. Treatment with tamoxifen was associated with worse 10-yr overall survival (OS) in women whose tumors overexpressed c-erb-B2. In contrast, women with c-erb-B2-negative disease who received tamoxifen had improved OS (Table 2) (28). In a recent update of this report, c-erb-B2 was one of eight markers that were retrospectively assayed in tissues from 83% of the patients enrolled in the original clinical trial (i.e., with or without chemotherapy). With a median follow-up of 15 yr, hazard ratio (HR) of death of tamoxifen over no-tamoxifen for c-erb-B2-positive subjects was 1.09 (95% confidence interval [CI]: 0.63–1.87, p = 0.04) compared to HR of 0.59 in women who did not overexpress c-erb-B2 (95% CI: 0.40-0.87) (29). The presence of c-erb-B2 was predictive of response to tamoxifen in the subgroup of patients with hormone receptor-positive tumors. The results of this updated report suggest that women with tumors overexpressing c-erb-B2 have a worse outcome in the presence of 2 yr of tamoxifen; however, this update included premenopausal patients who received adjuvant systemic chemotherapy with the CMF regimen (cyclophosphamide, methotrexate, 5-fluorouracil) in both groups. HR for c-erb-B2-negative patients who received only tamoxifen vs no systemic therapy was 0.54 (0.47-1.14) compared to 2.23 (0.95-5.23) for c-erb-B2-positive patients. These results were not statistically significant. When comparing women who received CMF and tamoxifen to those who received CMF only, significant differences in HR of response to tamoxifen were not seen. The multiple subgroup analyses of the GUN-1 study reflect the many shortcomings that are associated with investigations of the predictive role of c-*erb*-B2 in breast cancer.

In the Cancer and Leukemia Group B (CALGB) study 8541, 1572 women with node-positive breast cancer were randomly assigned to one of three dose groups of the CAF combination (cyclophosphamide, doxorubicin, 5-fluorouracil). In an analysis of 650 women with ER-positive disease who received 5 yr of adjuvant tamoxifen vs not in a nonrandomized fashion after the completion of CAF, tamoxifen was associated with similar risk reduction in DFS in the c-erb-B2-positive and c-erb-B2-negative groups (Table 2) (30). Although these results are from a nonrandomized assignment of tamoxifen, the number of samples available for the marker study was large, several methods assessing c-erb-B2 status were used, and the patient population was relatively homogeneous (node-positive, hormone receptorpositive). However, all the patients included in the analysis received doxorubicin-based therapy, albeit one of three-dose levels, prior to tamoxifen. Given the clinical data suggesting that c-erb-B2-positive tumors may be associated with relative sensitivity to anthracyclines (reviewed below), it can be hypothesized that doxorubicin may reverse relative resistance to tamoxifen. Other reports from nonrandomized investigations suggested a relative resistance to tamoxifen for the subgroups of patients with c-erb-B2positive breast cancer vs those with marker-negative disease (31-33).

Others suggested that the optimal treatment duration of tamoxifen might differ for women with c-*erb*-B2–positive vs –negative disease. c-*erb*-B2 status was assessed either by DNA amplification assay (n = 181) or by flow cytometry of the protein (n = 396) in women who received 2 or 5 yr of adjuvant tamoxifen in a randomized clinical trial. Patients with c-*erb*-B2–negative disease had a significant benefit from 5 yr of tamoxifen treatment (relative risk [RR] = 0.62, 95% CI: 0.42–0.93), while no added benefit was seen for c-*erb*-B2–positive patients (RR = 1.1, 95% CI: 0.41–3.2) (*34*). However, when the analysis was restricted to ER-positive patients, the difference in relative hazard was not statistically significant (p = 0.065).

Recently, results were reported from a large clinical trial of adjuvant oophorectomy and tamoxifen (20 mg/day for 5 yr) or observation. From 1993 to 1999, 709 premenopausal women with operable breast cancer were recruited to this trial. Hormone receptor status, which was not available at the time of recruitment, was subsequently evaluated in 66% of the participants, of those 62% were ER positive (i.e., 288 women or 41% of the study participants). With a median follow-up of 3.6 yr, 5-yr DFS and OS were improved for the women in the treatment group with hormone receptor–positive disease (*35*). Overall, 282 samples were available for IHC analysis. The HR for DFS in c-*erb*-B2–positive patients with adjuvant endocrine therapy was 0.37 (95% confidence interval [CI], 0.26–0.89) compared to 0.48 (95% CI, 0.31–0.71) for c-*erb*-B2–negative patients, suggesting that

both marker groups benefited from the treatment equally. Likewise, the HR for OS was similar for c-*erb*-B2–positive and –negative patients. When analyses were restricted to patients with ER-positive tumors, c-*erb*-B2 positivity was associated with worse DFS and OS compared to women with c-*erb*-B2–negative tumors. These results suggest that women with c-*erb*-B2–positive and –negative disease derive similar proportional benefit from a combination of oophorectomy and tamoxifen, however, it is possible that the c-*erb*-B2–positive tumors may be associated with a more aggressive biology (negative prognostic factor) and the outcomes, although improved with the treatment, may still be inferior to outcomes of women with c-*erb*-B2–negative disease.

2.4. c-erb-B2 as a Predictive Factor for Response to Endocrine Therapies: Conclusions

The results outlined above demonstrate the multiple limitations of the studies reported to date that evaluated the predictive role of c-*erb*-B2 to endocrine therapies. Most studies are very small, with patients who may or may not have received a uniform treatment, and may or may not have been enrolled in a single clinical trial. The assays used to determine c-*erb*-B2 status vary among different investigations, as do the antibodies used, cutoff levels, and scoring algorithms. While initial reports supported preclinical studies that suggested that c-*erb*-B2 may indeed be associated with a worse response to endocrine manipulations, recent reports indicate that women with either c-*erb*-B2–positive or –negative disease derive similar proportional benefit from endocrine treatments, and/or that not all endocrine treatments are alike.

Given the deficiencies of most marker studies and inconsistent results from larger randomized studies, it is difficult to conclude whether c-erb-B2 is predictive of response to all or to specific hormone therapies. Based on the available data, hormone therapies should not be withheld from women simply because their tumors overexpress or amplify c-erb-B2. Hormone treatment recommendations should be based on hormone receptor status, risk of relapse, menopausal status, and comorbidities. Until further data are available, it may be reasonable to offer adjuvant anthracycline-based chemotherapy in addition to tamoxifen to women with primary breast cancer with expression of hormone receptors and overexpression/amplification of c-erb-B2. Whether the addition of ovarian ablation to tamoxifen in pre- or perimenopausal women with small hormone receptor-positive c-erb-B2positive disease will reverse possible tamoxifen resistance is simply not known. Likewise, conclusive data to support the use of aromatase inhibitors instead of tamoxifen in postmenopausal women with c-erb-B2-positive disease are not available.

3. c-erb-B2 AS A PREDICTIVE FACTOR FOR RESPONSE TO CHEMOTHERAPY 3.1. Preclinical Data

Results from preclinical experiments suggested that breast cancer cell lines that overexpress or amplify the c-erb-B2 receptor have not provided conclusive evidence regarding c-erb-B2 status and relative sensitivity or resistance to chemotherapy. In breast cancer cell lines that were transfected with c-erb-B2, significant difference in the response to seven different chemotherapy agents was not observed between the wild-type or transfected cells (36). Similarly, normal human mammary epithelial cells that were designed to overexpress c-erb-B2 did not demonstrate resistance to singleagent doxorubicin, paclitaxel, cisplatin, 5-fluorouracil, or methotrexate (37). It is possible that overexpression of the oncogene by itself may not be sufficient to confer relative sensitivity or resistance to common chemotherapies. In human cancers c-erb-B2 positivity may be associated with several additional subcellular changes. For example, breast cancer specimens commonly coamplify c-erb-B2 and the enzyme topoisomerase II, the target of anthracyclines (38). Investigators thus evaluated c-erb-B2 status in 40 cell lines that were primarily derived from tissue of chemotherapynaïve patients at the time of surgery. The tumors were then exposed to six different concentrations of CEF (cyclophosphamide, epirubicin, 5-fluorouracil) or CMF (cyclophosphamide, methotrexate, 5-fluorouracil) combination. Tumors with an intermediate or strong overexpression of the marker were more sensitive to either CMF or CEF compared to tumors with low or no c-erb-B2 expression. Of note, the results were statistically significant when the antibodies TAB250 and AO485 were used (p values 0.044 and 0.032, respectively) but not with the CB11 antibody (p = 0.8) (39).

Other studies focused on response to taxanes. Breast cancer cell lines transfected with c-*erb*-B2 blocked paclitaxel- or docetaxel-induced apoptosis (40,41). Furthermore, infection of the taxane-resistant cells with adenovirus type 5 *EIA* led to down-regulation of c-*erb*-B2 and subsequent restoration of paclitaxel sensitivity (42).

3.2. CMF and Other Alkylating Agent-Based Therapy

Several uncontrolled clinical trials have demonstrated that patients who were treated with adjuvant CMF or CMF-containing regimens and whose tumors were c-*erb*-B2 positive had worse outcomes compared to patients with c-*erb*-B2–negative disease (31,43-45). In contrast, in the metastatic setting, patients with c-*erb*-B2–positive cancers were more likely to respond to CMF compared to those with c-*erb*-B2–negative tumors (17,46). Because of heterogeneity in patient population, small sample size and variability in methods of c-*erb*-B2 detection, a conclusive correlation between c-*erb*-B2

status and CMF cannot be made. Results from controlled studies demonstrated no correlation, relative resistance, or relative sensitivity to the CMF regimen (Table 3).

The first two controlled studies were reported more than a decade ago, and both suggested that c-erb-B2 may be a negative predictive factor for response to CMF-like regimens. Blocks were available from 306 nodenegative women enrolled in U.S. Intergroup 0011. The women received cyclophosphamide, methotrexate, 5-fluorouracil, and prednisone (CMFP) or no systemic therapy. DFS was improved for c-erb-B2-negative women who were on the treatment group compared to control (80% vs 58%, p =0.0003). However, c-erb-B2-positive women had similar DFS regardless of the treatment assignment (78% vs 68%, p = not significant) (47). In International Breast Cancer Study Group (Ludwig) Trial V, node-positive women received six cycles of CMF vs one preoperative cycle of CMF. Node-negative women received one perioperative cycle of CMF vs no therapy. Postmenopausal women also received tamoxifen. Marker analysis was performed in 60% of the clinical trial participants. In c-erb-B2-negative patients, six cycles of CMF were superior to one cycle of therapy. However, outcomes were similar for women with c-erb-B2-positive disease who received six cycles or one cycle of CMF (Table 3) (48).

More recently, British investigators compared outcomes of 274 nodepositive women who received six cycles of CMF vs no adjuvant systemic therapy. For women with c-*erb*-B2–negative tumors, median survival was improved with CMF compared to the control group (12.7 vs 7.3 yr, p =0.0014) (49). Women whose tumors were c-*erb*-B2 positive had a worse survival compared to c-*erb*-B2–negative patients regardless of treatment assignment. However, CMF therapy was associated with an almost 2-yr survival benefit for c-*erb*-B2–positive patients (median survival 6.1 and 4.4 yr for CMF-treated and control patients, respectively, p = 0.08). In a multivariate analysis, c-*erb*-B2 positivity was marginally associated with worse survival (p = 0.03); however, the c-*erb*-B2–positive patients were also more likely to have ER-negative tumors and four or more involved nodes.

In another large study, 386 node-positive women were randomly assigned to 12 cycles of CMF vs no adjuvant chemotherapy, and 337 samples were available for IHC of *c-erb-B2 (50)*. HR for DFS for women treated with CMF vs not was 0.484 for *c-erb-B2*–positive patients, compared to 0.641 for *c-erb-B2*–negative patients (Table 3). HR for cause specific survival for *c-erb-B2*–positive patients was 0.495 compared to 0.73 in *c-erb-B2*–negative patients, suggesting that women benefited equally from CMF regardless of *c-erb-B2* status.

Other small studies evaluated c-*erb*-B2 status and response to cyclophosphamide-based high-dose chemotherapy. In aggregate, the results suggest a worse DFS for women with tissue c-*erb*-B2 overexpression or high concentration of ECD–c-*erb*-B2. The patients included in these studies may have received a variety of different regimens prior to the high-dose therapy and the results may also be confounded by prior use of arthracyclines.

Taken together, results from large prospective clinical trials of CMF vs no therapy suggest that women with *c-erb*-B2–positive or –negative tumors derive benefit from the CMF regimen. However, women with *c-erb*-B2– positive disease may have a worse prognosis at baseline and thus even if the proportional benefit from CMF is equivalent in marker-positive or -negative patients, the marker-positive patients may still suffer worse outcome following CMF chemotherapy compared to women with *c-erb*-B2–negative tumors.

3.3. Anthracycline-Based Regimens

While several studies evaluated c-*erb*-B2 status and response to CMF vs no chemotherapy, most studies that correlated the role of the marker and response to anthracycline-based therapy included comparisons either to non-anthracyline–containing regimens or between different schedules and doses of the anthracycline (Table 3).

In SWOG Trial 9445 (Intergroup 0100), 1470 node-positive, ER-positive, postmenopausal women were randomly assigned to CAF combination with tamoxifen administered sequentially or concurrently (designated CAFT) vs tamoxifen alone. c-*erb*-B2 analysis was performed on 595 samples (41%) using IHC. CAFT was marginally superior to tamoxifen for the entire study population; however, women whose tumors overexpressed the c-*erb*-B2 receptor had substantial benefit form the addition of CAF to tamoxifen. Women whose tumors did not over express c-*erb*-B2 did not gain benefit from the addition of CAF to tamoxifen (51).

In National Surgical Adjuvant Breast and Bowel Project (NSABP) study B-11, node-positive women were randomly assigned to melphalan and 5-fluorouracil with or without doxorubicin (PF or PAF). With a 13.5-yr median follow-up, outcomes were improved for women who received the doxorubicin-based regimen. For c-*erb*-B2–positive patients, RR for DFS was 0.6 (0.44–0.83, p = 0.001), and for OS it was 0.66 (0.47–0.92, p = 0.01). For c-*erb*-B2–negative patients DFS and OS were not significantly different between the two treatment groups (52).

3.3.1. ANTHRACYCLINE-BASED VS CMF-LIKE REGIMENS

Based on early studies that suggested that women overexpressing c-*erb*-B2 may be relatively resistant to CMF-like regimens and sensitive to anthracyclines, studies that compared response to either of these regimens were of great interest (Table 3). In one study comparing CMF vs CAF, the 5-yr OS of the CMF-treated group was 84% for c-*erb*-B2–negative patients

Y	Randomized Clinical Trials of Adjuvant CMF-Based and Anthracycline-Based Regimens	I rials of Adjuvant (CMF-Based and An	thracycline-Based	Regimens	
	Number of patients in marker study		HR disease-free su (95% CI, p value) arm 1 vs arm 2	HR disease-free survival (95% CI, p value) arm 1 vs arm 2	HR overc (95% C1 arm I	HR overall survival (95% CI, p value) arm 1 vs arm 2
Author and study	(% c-erb-B2-positive)	Treatment arms	c-erb-B2-positive	c-erb-B2-positive c-erb-B2-negative	c-erb-B2–positive	c-erb-B2-negative
CMF-like Intergroup 0011 (47)	406 (14%)	 CMFP No therapy 	78% vs 68% ^a (NS)	80% vs 58% ^a (0.0003)	NR	NR
Ludwig V (node positive group) (48)	746 (19%)	1. CMF × 6 2. CMF × 1	0.77 (0.51–1.16)	0.57 (0.46–0.72)	46% vs 40% (0.93)	71% vs 61% (0.01)
Miles (Guy's Hospital) (49)	274 (30%)	1. CMF × 6 2. No therapy	NR	NR	6.1 vs 4.4 yr (0.08)	12.7 vs 7.3 yr (0.0014)
Menrad (Milan) (50)	337 (16%)	1. CMF × 12 2. No therapy	$\begin{array}{c} 0.484 \\ (0.284{-}0.827, \\ 0.004) \end{array}$	$\begin{array}{c} 0.641 \\ (0.481 - 0.853, \\ 0.001) \end{array}$	$\begin{array}{c} 0.495 \\ (0.287 - 0.853, \\ 0.006) \end{array}$	$\begin{array}{c} 0.73\\ (0.537-0.991,\\ 0.22)\end{array}$
Anthracycline-based therapy Anthracycline-based vs not	202 (12 02)		0 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	0 1 07 0 1 07 a	đ	đŅ
(IC) CHHADOMS	(2/01) (60	1. CALI 2. Tamoxifen	(0.01)	(0.39)	NN	NN
NSABP B-11 (52)	638 (37%)	1. PAF 2. PF	$\begin{array}{ccc} 0.6 & 0.96 \\ (0.44\text{-}0.83, 0.001) & (0.75\text{-}1.23, 0.74) \end{array}$	0.96 (0.75-1.23, 0.74)	0.66 (0.47–0.92, 0.01)	0.9 (0.69–1.19, 0.47)
Anthracycline vs CMF NSABP B-15 (86)	1355 (30%)	1. AC 2. CMF	0.84 (0.65–1.07, 0.15)	1.02 ($0.86-1.2, 0.84$)	0.82 (0.63–1.06, 0.14)	1.07 (0.88–1.3, 0.51)
Vera (53)	141 (13%)	1. CAF 2. CMF	NR NR	NR NR	72% vs 42% (0.3)	82% vs 84% (NS)

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DiLeo (54)	354 (21%)	1. HEC 2. EC 3. CMF	3 vs 1:1.42 (0.54–3.76, 0.48) 3 vs 2:1.65 (0.66–4.13, 0.29)	3 vs 1:0.84 (0.49–1.44, 0.53) 3 vs 2:0.66 (0.39–1.1, 0.11)	NR	NR
NCI-C MA.5 (56)	602 (24%)	1. CEF 2. CMF	1.54 (0.06)	1.07 (0.61)	1.29 (0.28)	0.90 (0.53)
Moliterni (57)	506 (19%)	1. CMF × 8 6 A × 4 2. CMF × 12	0.83 (0.46–1.49)	1.22 (0.91–1.64)	0.61 (0.32-1.16)	1.26 (0.89–1.79)
Anthracycline dose CALGB 8541 (58,59)	992 (24%)	1. High-dose CAF 71%, 52%, 51% ^a 65%, 66%, 60% ^a 2. Moderate-dose CAF (<0.01) (0.058) 3. Low-dose CAF	71%, 52%, 51% ^a F (<0.01)	$65\%, 66\%, 60\%^a$ (0.058)	87%, 66%, 63% ^a (<0.01)	77%, 82%, 78% ^a (0.048)
DiLeo (54)	354 (21%)	1. HEC 2. EC	2 vs 1:0.93 (0.31–2.77, 0.9)	2 vs 1:0.93 2 vs 1:1.33 (0.31–2.77, 0.9) (0.82–2.14, 0.25)	NR	NR
CAF, Cyclophosphamide, doxorubicin, 5-fluorouracil; CI, confidence interval; EC, epirubicin, cyclophosphamide; HEC, high-dose epirubicin,	doxorubicin, 5-fluo	rouracil; CI, confidenc	e interval; EC, ep	irubicin, cyclophos	sphamide; HEC, hi	gh-dose epirubicin,

, r 2 -ngm y vyvivpiuop. יישי ישם (at CAF, Cychopnospinatinue, uoxonuorent, 2-muoroutaeut, Ci, vomueure cyclophosphamide; HR, hazard ratio; NR, not reported; NS, not significant. #% DFS or OS per treatment arm.

compared to 42% in those who overexpressed the marker (p = 0.006) (53). In contrast, women who were treated with CAF had similar 5-yr OS regardless of c-*erb*-B2 status. These results were consistent with the hypothesis that women with c-*erb*-B2–positive disease were more likely to benefit from CAF while those with c-*erb*-B2–negative disease derived equal benefit from CMF or CAF.

European investigators evaluated c-erb-B2 status in 354 samples from 777 node-positive patients who were randomly assigned to six cycles of CMF vs eight cycles of either moderate dose epirubicin and cyclophosphamide combination (EC) or high-dose epirubicin and cyclophosphamide (HEC). Overall, HEC was associated with improved outcomes compared to EC, while the CMF and EC groups had equivalent outcomes. However, when outcomes were analyzed by c-erb-B2 status and treatment, markerpositive women were more likely to benefit from HEC or EC compared to CMF (Table 3) (54). A significant difference was not seen between EC and HEC in either c-erb-B2-positive or -negative patients. However, the sample size of each group was small, and all comparisons were not significant with wide and overlapping confidence intervals. In another study, 348 women received adjuvant CMF or weekly epirubicin and 266 samples were available for marker analysis. OS was worse for women with c-erb-B2-positive disease in both groups (p = 0.02), but a significant difference in OS was not observed between the arms (p = 0.12) (55).

In National Cancer Institute of Canada (NCI-C) study MA .5, 710 premenopausal node-positive women were randomly assigned CEF or CMF, and tissue for c-*erb*-B2 analysis was available from a large proportion (85%) of the study participants. DFS was statistically significantly worse for the patients whose tumors overexpressed or amplified c-*erb*-B2 (56). These results suggested that women with marker-positive disease may have preferential sensitivity to CEF; however, data were not conclusive owing to a small sample size and a short follow-up. In another recent report, specimens from 506 patients who were included in a prospective study of 12 cycles of CMF vs 8 cycles of CMF followed by 4 cycles of doxorubicin were analyzed for c-*erb*-B2 status. With a 15-yr median follow-up, recurrence-free survival and OS were improved for patients who overexpressed c-*erb*-B2 who received sequential CMF followed by doxorubicin compared to the CMFonly group (57).

3.3.2. ANTHRACYCLINE DOSE

Results from CLAGB study 8541 suggested that c-*erb*-B2 overexpression or amplification may be predictive for response to higher dose levels of doxorubicin. In this study, women received one of three dose levels of CAF, designated low, moderate, and high dose. DFS and OS were improved for women with c-*erb*-B2–positive tumors who received high-dose CAF (p < 0.01) (58). The results were confirmed with a larger data set and with other methods including fluorescence *in situ* hybridization (FISH) and polymerase chain reaction (PCR) (Table 4) (59,60). However, the regimen designated "high-dose" is now considered standard while the moderate- and low-dose regimens may be suboptimal alternatives. Whether a higher than standard dose of doxorubicin will be associated with even further benefit for c-*erb*-B2–positive women is not known. The European study comparing CMF to EC or HEC outlined earlier, did not demonstrate a preferential outcome with HEC compared to EC (54). Marker analysis from patients included in CALGB study 9344 who received one of three escalating doses of doxorubicin (60, 75, or 90 mg/m²) may provide additional information (61).

Although the studies described in the preceding are very heterogeneic, the data overall suggest that while both CMF and anthracyclines are effective treatments for c-*erb*-B2–positive or –negative breast cancer, it is possible that c-*erb*-B2–positive patients derive greater benefit from anthracycline-based therapy compared to CMF-like regimens. At the same time, c-*erb*-B2–negative patients may benefit equally from either type of therapy. While clinicians may choose anthracyclines preferentially for c-*erb*-B2–positive patients, CMF should be offered to women for whom anthracycline-based therapy is not appropriate regardless of c-*erb*-B2 status.

3.4. Taxane-Based Regimens

Preclinical studies suggested that c-*erb*-B2 overexpression may result in relative resistance to taxanes; however, reports from clinical trials are mixed. Taxanes have been available for more than a decade and only a handful of clinical studies evaluated the role of c-*erb*-B2 in predicting response to taxanes. Most studies were in the metastatic setting and included combination of taxanes and anthracyclines. In a study of women with metastatic breast cancer who received single agent paclitaxel (n = 106) or docetaxel (n = 20), the odds of responding to the taxane for c-*erb*-B2–positive patients were greater compared to the c-*erb*-B2–negative group. Women with c-*erb*-B2–positive disease had a response rate of 47% compared to a 39% response rate in the c-*erb*-B2–negative group (p = 0.027) (62,63). c-*erb*-B2 was associated with a relative sensitivity to either taxane.

Colomer and colleagues evaluated the predictive role of tissue (IHC) or circulating ECD–c-*erb*-B2 and response to a combination of doxorubicin and paclitaxel administered as first-line chemotherapy to women with meta-static breast cancer. With a 23-mo follow-up, the authors reported that elevated circulating ECD–c-*erb*-B2 correlated with worse response rate (complete response 0% vs 26% for c-*erb*-B2–positive and –negative patients, respectively, p = 0.35) (64). However, response to the therapy was similar for patients with tissue c-*erb*-B2–positive or –negative disease. Importantly, circulating ECD–c-*erb*-B2 correlated with bulk of disease and

		Randomiz	zed Clinical Tri	als of Taxane-Co	Randomized Clinical Trials of Taxane-Containing Regimens vs Not	iens vs Not		
	Patient population and number in marker study		Response rate (p value)	tte (p value)	Median time to pro arm 1 v	Median time to progression (p value) Median overall survival (p value) arm 1 vs arm 2	Median overall s arm I v	verall survival (p value) arm 1 vs arm 2
Author	(% c-erb-B2-positive) Treatment arms c-erb-B2-positive c-erb-B2-negative c-erb-B2-positive c-erb-B2-negative c-erb-B2-negative	Treatment arms	c-erb-B2-positive	c-erb-B2-negative	c-erb-B2-positive	c-erb-B2-negative	c-erb-B2-positive	c-erb-B2-negative
Konecny (87)	Metastatic 256 (38%)	1. ET 2. EC	76% vs 46% (0.04)	50% vs 33% (0.02)	10.5 vs 7.1 mo (0.116)	9.6 vs 10.4 mo (0.35)	21.4 vs 16.4 mo 27.5 vs 33.1 mo (0.319) = (0.292)	27.5 vs 33.1 mo = (0.292)
Di Leo (66)	Metastatic 176 (20%)	 Docetaxel Doxorubicin 	67% vs 27% (0.04)	40% vs 35% (0.70)	7 vs 4.7 mo (0.73)	5 vs 5.9 mo (0.21)	14.4 vs 10.8 (0.32)	12.6 vs 16.9 (0.07)
Nabholtz (68)	Adjuvant 1491 (20%)	1. TAC 2. FAC	AN	NA	HR: 0.59 (95% CI 0.38– 0.91, 0.02)	HR: 0.74 (95% CI 0.54– 1.01, 0.06)	NR	NR
CI, Confi HR, hazard r	CI, Confidence interval; EC, epirubicin and cyclophosphamide; ET, epirubicin and paclitaxel; FAC, 5-fluorouracil, doxorubicin, cyclophosphamide; HR, hazard ratio; NA, not applicable; NR, not reported; TAC, docetaxel, doxorubicin, cyclophosphamide.	epirubicin and cy ole; NR, not repor	/clophosphamide rted; TAC, doceta	; ET, epirubicin ; ixel, doxorubicin,	and paclitaxel; F _i cyclophosphamic	AC, 5-fluorouracil le.	l, doxorubicin, cy	clophosphamide;

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may be a more quantitative assay compared to IHC. In a similar study of 49 patients who received combination doxorubicin and paclitaxel, overall response was equivalent in women with c-*erb*-B2–positive or –negative disease (90% and 93%, respectively). However, the percentage of women free from progression at 20 mo was greater in the c-*erb*-B2–positive group (63% and 48%, respectively) (65).

In the Eastern Cooperative Oncology Group Study (ECOG) 1193, 107 women with metastatic breast cancer received combination of doxorubicin and paclitaxel, paclitaxel alone (n = 109), or doxorubicin alone (n = 64). The group who received doxorubicin alone was then allowed to cross over to paclitaxel. The authors correlated response to taxane (alone or in combination) with c-*erb*-B2 status. A difference in response was not observed among the c-*erb*-B2–positive and –negative patients (p = 0.51). However, women with c-*erb*-B2–positive disease had a worse survival compared to the c-*erb*-B2–negative group (p = 0.0008).

In a study of 326 women with metastatic disease were randomly assigned to single-agent doxorubicin or docetaxel, samples for c-*erb*-B2 analysis by FISH were available from 176 participants. Objective response, TTP, and OS were similar for women in either treatment group. However, when analyzed by c-*erb*-B2 status, docetaxel provided greater benefit compared to doxorubicin in the marker-positive group (Table 4) (*66*). Marker-negative women had similar outcomes with either treatment.

Two other studies compared anthracycline-containing regimens with or without docetaxel. Konecny compared outcomes of women with metastatic breast cancer who were treated with combination epirubicin and cyclophosphamide (EC) or epirubicin and docetaxel (ET). ET-treated women with c-*erb*-B2-amplified tumors had improved response rate, DFS and OS as compared to those who received EC (Table 4) (67). However, women whose tumors did not amplify the oncogene had similar response to either regimen. Others compared adjuvant combination FAC and TAC in women with node-positive breast cancer. While TAC was superior to FAC in all women, c-*erb*-B2–positive women derived substantial benefit from TAC (HR 0.59, p = 0.02), while the benefit for marker-negative patients was approaching but not statistically significant (HR 0.74, p = 0.06) (68).

3.5. c-erb-B2 as a Predictive Factor for Response to Chemotherapy: Conclusions

Review of randomized clinical trials suggests that c-*erb*-B2 may indeed be predictive of response to specific chemotherapies. It is also possible that c-*erb*-B2 is a surrogate marker that reflects a more aggressive tumor biology and correlates with other factors that may predict for response to chemotherapy (e.g., lack of hormone receptors and worse grade). Based on the available data, systemic chemotherapy recommendations should be made based on the patient's risk of recurrence using accepted markers such as stage, nodal status, tumor size, grade, and hormone receptor status. The data overall suggest that women with c-*erb*-B2–positive disease may be more sensitive to anthracyclines and if a contraindication does not exist may be the combination of choice for those women. However, CMF should be considered for women who cannot tolerate anthracyclines. Sufficient evidence to preferentially recommend combination of anthracycline and taxane to women with c-*erb*-B2–positive disease is not available. Likewise, there is not sufficient evidence to recommend escalation of standard dose of anthracyclines to women with c-*erb*-B2–positive tumors.

4. c-*erb*-B2 AS A PREDICTIVE FACTOR FOR RESPONSE TO TRASTUZUMAB

Trastuzumab (Herceptin) is a humanized monoclonal antibody that targets the extracellular domain of c-*erb*-B2. In preclinical trials, trastuzumab suppressed tumor growth in breast cancer xenografts. Combination of trastuzumab with several chemotherapy agents such as doxorubicin, paclitaxel, and cisplatin provided synergistic tumor suppression (69).

The efficacy of trastuzumab has been examined in several clinical trials in metastatic breast cancer. In women with metastatic breast cancer whose tumors overexpressed c-*erb*-B2, single-agent trastuzumab was associated with a response rate of 12–15% in heavily pretreated women and 26% as first-line therapy (70–72). The initial clinical trials of single-agent trastuzumab have included patients with 2+ or 3+ overexpression of c-*erb*-B2 using IHC. In a study that evaluated response to single-agent trastuzumab in 111 women, response rate was 35% in those with a 3+ expression and 0% in women with a 2+ expression. When the same patients were evaluated using FISH, response rate in patients who amplified the oncogene was similar to those with the 3+ expression (72).

Based on the encouraging results of studies of single-agent trastuzumab, several uncontrolled and controlled studies evaluated combination of this agent and chemotherapy. In the largest randomized clinical trial, 469 women received first-line chemotherapy with or without trastuzumab. Women who received prior anthracyclines received paclitaxel with or without trastuzumab, while women who had not received anthracyclines previously were treated with combination doxorubicin and cyclophosphamide with or without trastuzumab. Objective response (50% vs 32%, p < 0.001) and TTP (7.4 vs 4.6, p < 0.001) were improved in women who received chemotherapy and trastuzumab compared to chemotherapy alone (73). More importantly, 1-yr survival was significantly improved for women who received trastuzumab (79% vs 68%, p < 0.01). Owing to the high rate of cardiac toxicity seen on the trastuzumab and doxorubicin/cyclophosphamide arm, the combination was not recommended for clinical use.

In other studies, combination trastuzumab and chemotherapy agents such as cisplatin, docetaxel, gemcitabine, paclitaxel, vinrelbine, or combination carboplatin and paclitaxel provides a higher response rates than expected with either agent alone (74–79). Most studies included only patients with overexpression or amplification of c-*erb*-B2. In one study that enrolled patients regardless of c-*erb*-B2 status, combination of paclitaxel and trastuzumab provided improved response rates compared to paclitaxel alone in women who overexpressed c-*erb*-B2 (81% and 67%), while no difference in response was observed in women without c-*erb*-B2 expressing tumors (46% and 41%) (77). Indeed, this study demonstrated that c-*erb*-B2 is the target of trastuzumab and c-*erb*-B2 status should be used for patient selection.

Studies of trastuzumab in the preoperative setting with paclitaxel or docetaxel demonstrated high response rates of 75% (78,80). MD Anderson Cancer Center investigators have recently demonstrated that the addition of trastuzumab to preoperative anthracycline and taxane-based chemotherapy was associated with a marked increase in pathological complete response (81). The role of adjuvant trastuzumab with or following systemic chemotherapy is under extensive investigation. Recent results from large cooperative group clinical trials designed to evaluate the efficacy and toxicity of trastuzumab in the adjuvant setting were recently presented in abstract form. The addition of trastuzumab to standard adjuvant systemic therapy provided a substantial improvement in DFS and OS. Importantly, investigators have found discrepancies in evaluation of c-erb-B2 testing of patients enrolled in adjuvant randomized clinical trials of chemotherapy with or without trastuzumab compared to a central laboratory (82,83). Based on these evaluations, most centers and laboratories accept 3+ as overexpression but 2+ or 1+ is referred for further FISH testing. Other studies are evaluating the role of trastuzumab with other novel agents. Finally, several studies focus on newer agents and vaccines that target the c-erb-B2.

5. CONCLUSIONS

c-*erb*-B2 is clearly an important oncogene in breast cancer. However, after two decades of investigations, the predictive role of c-*erb*-B2 to specific treatments is still uncertain. With the exception of trastuzumab, c-*erb*-B2 status cannot be used to determine systemic treatment recommendations for breast cancer. Although some evidence suggests that women with c-*erb*-B2–positive disease may be less likely to benefit from a tamoxifen and CMF-like regimen, the data are mixed. Most controlled studies suggest that women with c-*erb*-B2–positive disease may have a preferential response to anthracycline-containing regimens. Thus, anthracycline-based therapy should be recommended to women with c-*erb*-B2–positive breast cancer unless there is a contraindication to the administration of this group of

agents (such as cardiac disease or prior anthracycline use). Current clinical data also suggest a possible sensitivity to taxane-based therapy, especially when combined with an anthracycline. Finally, trastuzumab-based therapy should be clearly considered as first-line therapy for women with c-*erb*-B2 disease who are hormone receptor negative, or those with hormone receptor–positive disease who have progressed on endocrine treatments. In the adjuvant setting, the addition of trastuzumab should be considered for women with high-risk disease.

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9

erb-B2 as a Therapeutic Target

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CONTENTS

INTRODUCTION THE DEVELOPMENT OF HERCEPTIN[®] HER2 TESTING AND THE IMPACT ON TREATMENT EFFECT FROM HERCEPTIN CONCLUSIONS REFERENCES

SUMMARY

Trastuzumab is a humanized monoclonal antibody that targets the type 1 tyrosine kinase receptor erb-B2 and has demonstrated survival benefit when used in combination with chemotherapy in the treatment of metastatic, HER2 overexpressing breast cancer. This chapter reviews the history of the development of trastuzumab including the biology of HER signaling, the technical issues of antibody "humanization," the diagnostic challenges in developing "targeted" therapeutic agents, and the clinical data that led to the approval of this agent for patients with HER2 overexpressing breast cancer.

Key Words: Breast cancer; HER2; erb-B2; trastuzumab; monoclonal antibody; targeted therapy.

1. INTRODUCTION

The human epidermal growth factor receptor 2 (HER2, c-erb-B2, HER2/ *neu*) is one of four known members of the type 1 family of tyrosine kinase

Cancer Drug Discovery and Development: Biomarkers in Breast Cancer: Molecular Diagnostics for Predicting and Monitoring Therapeutic Effect Edited by: G. Gasparini and D. F. Hayes © Humana Press Inc., Totowa, NJ receptors present in human epithelial cells. These receptors all share significant homology, structurally consisting of an extracellular (ligand binding) domain, a transmembrane region, and an intracellular domain containing a tyrosine kinase region (1). In addition to the 4 known receptors, 12 ligands have been described within this signaling pathway, each exhibiting highly specific binding to the extracellular domain of one or more of the receptors. More primitive species have a less diverse and robust HER signaling network. *C. elegans* contains a single receptor and ligand while *Drosophila* contains four ligands and a single receptor. This evolutionary richness and diversity is directly related to the critical role this signaling pathway plays in regulating multiple cellular functions including proliferation, differentiation, adhesion, migration, and survival.

During the past 25 yr much has been learned regarding the precise mechanism by which these various receptors and their ligands associate to initiate intracellular signal transduction pathways. Two of the receptors have peculiar characteristics. HER2 lacks any known ligand while HER3 has a "dead" kinase domain. Neither of these receptors can signal without associating or dimerizing with another member of the HER receptor family. Although weak signaling may occur with ligand binding to a single receptor (HER1 or HER4), receptor aggregation forming homo- or heterodimeric pairs or larger oligomeric structures result in sustained and potent signaling. In a series of elegant experiments, Yarden and colleagues demonstrated that heterodimers provide a significant increase in signaling potency over homodimers. In addition, they showed that heterodimers containing HER2 were the most potent signaling complexes, suggesting a central, pivotal role for HER2 in this pathway (2).

Recently the crystalline structures of HER1, in both its unbound and ligand bound conformations, HER2 and HER3 have been described (3–5). These structural observations show that receptor association occurs via a short hairpin loop within the extracellular domain II region of the receptors. HER1, in its non-ligand–bound state, assumes a "closed" conformation with this hairpin loop "buried" within a pocket of domain IV, When ligand binding occurs at a point between domains I and III there is a substantial structural change with pivoting of the receptor complex, disruption of the domain II–IV interaction, and exposure of the hairpin loop to form an "open" conformation available for receptor interactions. The structure of HER2 demonstrated that this critical hairpin loop is constitutively exposed, explaining the lack of need for a HER2 ligand. By adopting this fixed, "open" conformation, HER2 is always "ready to partner" with other receptor members and serves to explain the pivotal role that HER2 plays in this signaling pathway.

2. THE DEVELOPMENT OF HERCEPTIN®

Although these reports were published well after the development of Herceptin, earlier observations suggested that HER2 was an attractive target for therapeutic cancer drug development. Human HER2 was initially cloned by Ullrich and colleagues at Genentech in the mid-1980s and soon thereafter shown to be an oncogene (6). When *HER2* was transfected into normal cells they became transformed and displayed all the characteristics of malignant cells when grown in soft agar (Fig. 1) (7,8). At the same time, the initial reports were published suggesting that 25–30% of human breast cancers demonstrated an abnormality of HER2, specifically high-level amplification of a nonmutated copy of the gene (9). In this seminal observation, HER2 was found to be amplified in a subset of breast cancers and that amplification was shown to be a significant negative prognostic indicator, implying an integral role for HER2 in the biology of these cancers. This finding has been confirmed in many subsequent studies and there is considerable evidence that amplified HER2 is an early and stable molecular abnormality in breast cancer and an adverse prognostic indicator in both node-positive and node-negative disease (10-13). Because amplification is not associated with mutations or splice variants, corresponding increases in mRNA and protein expression by Southern, Northern, and Western blotting and immunohistochemistry were also seen (14). The observation that HER2 amplification was associated with high-level overexpression of its protein product, p185^{HER2}, provided a framework to consider strategies to antagonize or suppress this abnormal biologic phenotype in human breast cancer.

One strategic approach to suppress the function of a surface expressed protein product of an oncogene is the use of targeted monoclonal antibodies. In the late 1980s, a large (>100) library of murine antihuman HER2 antibodies was generated by immunizing mice with homogenates of human HER2 overexpressing cell lines (15,16). These antibodies were then screened to determine their specificity and binding affinity to HER2. A short list of antibodies with the highest K_d is shown in Table 1.

Following this initial screen, the most promising antibodies were evaluated for antitumor activity in both in vitro and in vivo model systems across a range of tumor cell lines. The antibody that was selected from this screening process for clinical development was known as 4D5. These preclinical studies also provided a critical observation that guided the overall clinical development plan for Herceptin.

From the data shown in Fig. 2, it was apparent that the antitumor activity of 4D5 was confined to cell lines that expressed moderate to high levels of HER2 (17). Although not unexpected, it did suggest that optimal clinical development of this antibody would require the parallel development of an assay to detect amplification or protein overexpression of HER2 in clinical

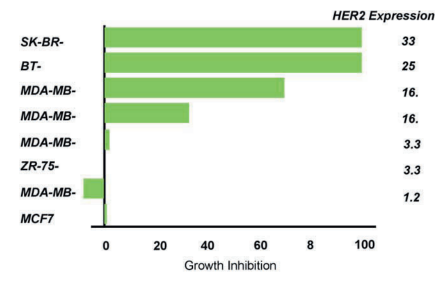


Fig. 1. Antibody inhibition of anchorage-independent growth is related to HER2 expression level.

ionocionar / meiooc	
$Kd^{ extsf{w}}\left(nm ight)$	Internalization in SK-BR-3
9.3	243
2.6	24
5.0	236
8.4	133
8.6	313
63	1801
	<i>Kd</i> [®] (<i>nm</i>) 9.3 2.6 5.0 8.4 8.6

Table 1 Binding Affinity and Internal Characteristics of Several Monoclonal Antibodies Against HER2

tumor material and then restrict enrollment in clinical trials to only those patients whose tumor demonstrated high levels of expressed HER2.

To standardize and validate an immunohistochemistry (IHC) assay to detect expressed p185^{HER2}, three breast cell lines were obtained from the American Type Culture Collection (ATCC); MDA-231, MDA-175, and SK-BR-3. The *HER2* gene copy number, using DNA slot blots (and subsequently by FISH to correct for chromosome 17 ploidy) and quantification of expressed HER2 receptors by Scatchard analysis, was determined as shown in Table 2. These cell lines served as performance controls for the development of a semiquantitative IHC assay, known as the Clinical Trials

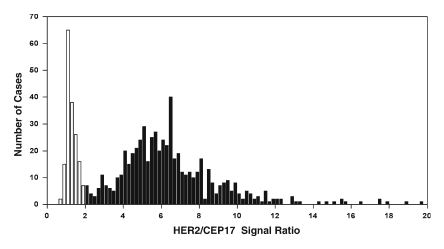


Fig. 2. Distribution of HER2/CEP17 signal ratios (n = 765).

Table 2
HER Expression Is Related to HER2 Amplification
in Cultured Human Breast Cancer Cell Lines

Cell line	HER2 receptors/cell	HER2/neu/CEP17 ratio
MDA-231	21,600	~1.1
MDA-175	92,400	~1.3
SK-BR-3	2,390,000	~4.5

Table 3
Immunohistochemistry Staining of Cultured
Human Breast Cancer Cell Lines to Develop IHC Score for HER2

Cell line	IHC score	Description
MDA-231	0	No discernible membrane staining
MDA-175	1+	Faint/barely perceptible, incomplete membrane staining
SK-BR-3	3+	Moderate/strong complete membrane staining

Assay (CTA) subsequently utilized to select patients for the Herceptin development trials.

4D5 was the antibody selected for the assay with a standard antimouse antibody and an avidin–biotin horseradish peroxidase complex. The antigen retrieval consisted of limited protease treatment. The conditions for the assay were standardized to obtain a reliable and characteristic membranestaining pattern from each of the performance control cell lines as shown in Table 3. After the development and validation of this assay, human clinical trials using antibody 4D5 (murine) were initiated. Only patients whose tumor scored 2+ or 3+ using the CTA were enrolled in all subsequent clinical trials. A total of 26 women with advanced metastatic breast cancer, known to be HER2 positive, were treated in a single dose and a multiple dose phase I trial with this murine antibody. Infusion reactions were common and both trials demonstrated the rapid and frequent development of neutralizing human antimouse antibodies (HAMA). Despite these findings, several patients in these trials demonstrated clear, but transient, objective tumor regression, establishing the "proof of concept" that antibodies to HER2 could result in antitumor effects in humans.

Advances in protein and antibody engineering in the late 1980s made it possible to conceptualize the idea of "grafting" the backbone of a human immunoglobulin molecule onto the consensus determining regions (CDRs) or binding domains of a specific murine monoclonal antibody; essentially "humanizing" the antibody. This concept would theoretically enable chronic, long-term dosing by avoiding the development of HAMAs and, with an intact and functional human F_c domain, could provide for additional antitumor activity by initiating targeted antibody-dependent cellular cytotoxicity (ADCC). Through a series of landmark technical achievements, a human IgG_1 molecule was successfully grafted onto antibody 4D5 (18). Hundreds of clones were created in this process to determine the precise sequence location where binding affinity began to degrade. The optimal clone was composed of 95% human and 5% murine sequences and was determined to have higher binding affinity to human HER2 and better antitumor activity in human xenograft model systems when compared with the native murine antibody, 4D5. This humanized antibody is now known as trastuzumab or Herceptin.

A small phase I program to establish the safety and pharmacokinetics of Herceptin was initiated in 1992. Infusion reactions continued to be seen with the humanized antibody in 30–40% of patients; however, they were rarely severe and typically confined only to the first infusion. The half-life was estimated to be 5–6 d and, most importantly, no HAMAs were detected with the humanized antibody. During the phase I period, additional preclinical studies were conducted to assess the interaction between Herceptin and several chemotherapeutic agents. Striking synergy was noted when HER2 overexpressing cells were exposed to the combination of Herceptin and the DNA damaging agent cisplatin (19). These observations led to the development of two phase II trials with Herceptin. The first trial (H0551g) evaluated single-agent Herceptin at the target dose of 4 mg/kg as a loading dose and then 2 mg/kg/wk until disease progression in highly refractory HER2-positive metastatic breast cancer. This trial showed that the drug was well tolerated and demonstrated an objective response rate of 11% (20). The second

Synergistic (CI <1)		Additive (CI = 1)	Subadditive (CI >1)
Vinorelbine	0.34	Doxorubicin	0.02 1110	Fluorouracil 2.87
Docetaxel/carboplatin	0.34	Paclitaxel	0.91	
Docetaxel	0.41	Epirubicin	0.99	
Etoposide	0.54	Vinblastine	1.09	
Cyclophosphamide	0.57	Methotrexate	1.36	
Paclitaxel/carboplatin	0.64			
Thiotepa	0.67			
Cisplatin	0.67			
Liposomal doxorubicin	0.7			
Gemcitabine	1.25–5.34 (*	variable, dose-o	dependent)	

Table 4 Trastuzumab and Chemotherapy: In Vitro Cytotoxicity Against HER2-Positive Breast Cancer Cell Lines^a

^{*a*}Based on a Combination Index (CI) score from multiple drug-effect analysis at fixed molar ratios. Pegram et al. *Oncogene* 1999;18:2241; Pegram et al. *Semin Oncol* 2000;27(Suppl 11):21; Slamon and Pegram. *Semin Oncol* 2001;28(Suppl 3):13.

trial (H0552g) attempted to exploit the preclinical observations of synergy by utilizing the novel combination of Herceptin and cisplatin. The eligibility for this trial required patients to have particularly aggressive disease, demonstrating progressive disease *during* treatment with prior chemotherapy. Despite selecting for this very poor prognostic feature, the trial demonstrated an overall response rate of 24% with an additional 24% of patients achieving either a minor response or stable disease (21). Subsequently a larger panel of agents was tested, in multiple HER2 overexpressing models, to guide the optimal combination of Herceptin with chemotherapeutic drugs (22). Table 4 shows the results of those experiments. Many agents demonstrate additive or synergistic activity, with vinorelbine, the platinum salts cisplatin and carboplatin, docetaxel, and the antiestrogen tamoxifen showing the greatest synergistic interactions. Doxorubicin, cyclophosphamide, and paclitaxel all showed additive interactions and 5-fluorouracil (5-FU) was the only agent tested that demonstrated less than an additive interaction.

The pivotal development program for Herceptin included two large clinical trials: one phase II trial evaluating the agent as monotherapy in relapsed breast cancer following one or two prior regimens for metastatic disease and a second phase III trial evaluating doxorubicin and cyclophosphamide (AC) with or without Herceptin in first-line therapy of metastatic disease. Patients in both trials were required to have HER2 overexpression with a score of 2+ or 3+ using the CTA performed at a single reference laboratory. Because of the increasing use of doxorubicin in the adjuvant setting and the approval of paclitaxel for the treatment of metastatic breast cancer, the phase III trial was amended approximately halfway through recruitment to provide for two different chemotherapy strata. Patients naïve to anthracyclines in the adjuvant setting continued to be treated with AC ± Herceptin while patients who had received anthracyclines in the adjuvant setting were assigned to paclitaxel \pm Herceptin. This second strata had more unfavorable baseline prognostic characteristics consisting of larger size tumors, higher lymph node burden, and a shorter disease-free interval following adjuvant chemotherapy. The phase III trial specified a minimum of six cycles of chemotherapy (AC or paclitaxel) with further chemotherapy at the discretion of the investigator. Few patients received more than six cycles of cytotoxic chemotherapy. Herceptin was given as a weekly dose concomitantly with chemotherapy and then continued until the point of disease progression, which was the primary end point of the clinical trial. At progression, patients in the control arm were eligible to receive Herceptin as part of a separate protocol and approx 65% went on to receive the drug from this treatment arm. Patients in the experimental arm were also permitted to continue to receive Herceptin after initial disease progression and approx 50% continued treatment in this arm. This aspect of the study design is important when assessing the results, given the possibility that this prespecified "crossover" design may have reduced some of the treatment effect from Herceptin on end points occurring after progression (i.e., overall survival).

The phase III trial (study H0648g) reached the predefined number of events in late 1997 and was unblinded, revealing a highly significant improvement in the primary end point, time to disease progression. The addition of Herceptin to standard chemotherapy for HER2 overexpressing metastatic breast cancer improved the time to disease progression from 4.6 to 7.4 mo (p = 0.0001; *see* Table 5). This treatment effect was seen in both chemotherapy strata (AC and paclitaxel), and secondary efficacy end points of response rate, duration of response, and time to treatment failure were also significantly improved with the addition of Herceptin to standard chemotherapy.

Although the trial was unblinded 9 mo after the last patient was enrolled, patients continued to be followed for survival until the final study analysis was completed 30 mo after the last patient was enrolled. This ensured that all patients were followed beyond the median survival time point. Despite the fact that 65% of the control patients crossed over and received Herceptin at the time of disease progression, median survival improved from 20.3 mo to 25.1 mo (p = 0.046) (23).

The large phase II trial (study H0649g) of single-agent Herceptin in HER2 overexpressing metastatic breast cancer following one or two prior chemotherapy regimens for metastatic disease also confirmed the clinical activity of Herceptin. A total of 222 patients were treated with Herceptin as a weekly dose and an overall response rate of 14% was confirmed by a blinded, external response evaluation committee (24). A subsequent clini-

			Efficacy (all en	Efficacy (all enrolled patients)	-	.
	Either chemotherapy	iotherapy	doxorubicin and c	doxorubicin and cyclophosphamide	Paclitaxel	xel
	+ $traztuzumab$ ($n = 235$)	Alone (n = 234)	+ traztuzumab $(n = 143)$	<i>Alone</i> (n = 138)	+ $traztuzumab$ ($n = 92$)	Alone (n = 96)
Complete response (CR) Partial response (PR)	18/235 (8) 96/235 (41)	11/234 (5) 66/234 (28)	12/143 (8) 63/143 (44)	9/143 (7) 50/138 (36)	6/92 (7) 33/92 (36)	2/96 (2) 13/96 (14)
Overall CR + PR 95%	114/235 (49) p < 0.001	74/234 (32) 001	75/143 (52)	$\begin{array}{ll} 2) & 59/138 \ (43) \\ p = 0.10 \end{array}$	39/92 (42) 15 p = <0.001	15/96 (16))01
Median response duration (mo)	9.3 <i>p</i> <0.001	5.9 001	9.1 $p = 0$	p = 0.0025	11.0 $p < 0.001$	4.4 01
Median time to disease progression (mo)	7.4	4.6	8.1	6.1	6.9	3.0
Relative risk (95% CI)	$\begin{array}{c} 0.51 \; (0.41, 0.63) \\ p < 0.001 \end{array}$	1, 0.63 001	0.62 (0.47, 0 <i>p</i> <0.001	$\begin{array}{l} 0.62 \; (0.47, 0.81) \\ p < 0.001 \end{array}$	0.38 (0.27, 0.53) p < 0.001	0.53) 01
Median time to treatment failure (mo)	6.9	4.5	7.2	5.6	5.8	2.9
Relative risk (95% CI)	0.58 (0.47, 0.70) p < 0.001	7, 0.70) 001	0.67 (0.52, 0.8) p = 0.001	0.67 (0.52, 0.86) p = 0.001	0.46 (0.33, 0.63) p < 0.001	0.63) 01
Median survival (mo) Relative risk (95% CI)	$25.1 \qquad 25.1 \qquad 2 \\ 0.80 (0.64, 1.00) \\ 0.46 \qquad -0.046$	20.3 4, 1.00) 046	26.8 0.82 (.6	8 21.4 0.82 (.61, 1.09) 5 - 0.16	$22.1 18 \\ 0.80 \ (.56, 1.11) \\ 0.18 \\ 0.18 $	18.4 1.11) 8
p = 0.00 Duration of response was defined as the time from first response to disease progression or death. Time to treatment failure was defined as the time from	P = 0.	response to disease	P = P.	Time to treatment fai	P = 0.1	he time from

Trastuzumah Efficacy in Prospective Randomized Clinical Trial of Datients with Metastratic Breast Cancer Table 5

randomization to disease progression, study discontinuation for any reason, use of additional anti-tumor therapy, or death. The final analysis of the primary end point was performed 9 mo after enrollment of the last patient. The most recent survival data were obtained 30 mo after enrollment of the last patient, with a median foilow-up of 35 mo (range: 30–51 mo). As of the data cutoff date of December 31, 1997, 388 of 469 (83%) of patients had discontinued the study, including 173 of 235 (74%) of patients receiving rhuMAb HER2 plus chemotherapy and 215 of 234 (92%) of patients receiving chemotherapy alone. From ref. 23 with permission. cal trial of single-agent Herceptin in previously untreated patients with HER2 overexpressing metastatic breast cancer (study H0650g) demonstrated an objective response rate of 26% (25). The three single-agent Herceptin trials (H0551g, H0649g, and H0650g) treated patients with highly refractory metastatic disease, refractory metastatic disease, and untreated metastatic disease, respectively, and demonstrated response rates of 11%, 14%, and 26%, suggesting that Herceptin is most effective when used early in the treatment of HER2-positive breast cancer.

3. HER2 TESTING AND THE IMPACT ON TREATMENT EFFECT FROM HERCEPTIN

All of the pivotal trials described in the previous section utilized an IHC assay known as the Clinical Trials Assay (CTA) to define eligible patients. Scores of 2+ or 3+ were defined as "positive," rendering the patient eligible for enrollment while scores of 0 or 1+ were defined as "negative." Although the pivotal trials were not adequately powered to evaluate definitively the impact of IHC score (2+ vs 3+) on treatment effect, the retrospective analysis did suggest that the treatment effect was seen predominantly in the patients whose tumors had the highest (3+) expression of HER2. In the single-agent trial, H0649g, the response rate was 6% in the 2+ group and 18% in the 3+ group. In the pivotal chemotherapy trial, H0648g, median time to disease progression in the 2+ group went from 5.6 to 6.6 mo while in the 3+ group the improvement was from 4.6 to 8.5 mo.

Further confusion around the definition of HER2 overexpression in the Herceptin pivotal trials was introduced with the development of the HercepTestTM assay. Although the CTA was used to select patients during clinical development, this assay was complicated and not well suited to commercialization. The HercepTest is a different IHC assay that was designed and developed to provide "similar" results to those provided by the CTA. In a large concordance study performed on more than 600 clinical specimens, the HercepTest assay was found to provide the same result as the CTA in 82% of individual samples. Although HercepTest was approved by the US Food and Drug Administration (FDA) to aid in the selection of patients for Herceptin therapy, there were no prospective or retrospective analyses of the predictive value of a score of 2+ or 3+ on the treatment effect of Herceptin. Unfortunately, tissue blocks were not archived from any of the pivotal Herceptin studies. Only tissue sections, 4–6 µm thick and mounted on glass slides, were retained from the patients' original tumor specimens. These type of specimens are subject to oxidation and other factors, resulting in significant antigen degradation over time and an unacceptable rate of "false-negative" IHC assay results. A small, pilot study of the HercepTest assay on archived material from the pivotal trials confirmed this high rate

of false-negative results and served to highlight the need to archive tissue blocks or tissue microarrays from patients enrolled in clinical trials of targeted agents such that retrospective analyses of various diagnostic assays can be performed and correlated with clinical results.

Since the introduction of the HercepTest assay for the selection of patients for Herceptin therapy, there has been increasing concern over the reliability of IHC assays to assess HER2 status accurately in routine (formalin-fixed and paraffin-embedded, or FFPE) clinical specimens. IHC assays for HER2 are subject to a number of potential shortcomings, including variability in fixation techniques, reagents, antigen retrieval methods, and subjectivity of staining interpretation. These shortcomings can result in both false-positive and false-negative results (26-30). Because the primary molecular abnormality resulting in HER2 overexpression in 25-30% of human breast cancer is amplification of a nonmutated copy of the gene, alternative diagnostic strategies to directly assess gene copy number were evaluated in the belief that these technologies might provide more consistent and precise information as compared to IHC. Fluorescence in situ hybridization (FISH) is a technique in which a highly specific complementary DNA sequence (probe) is developed to the gene of interest and then labeled (either directly or indirectly) with a florescent tag for eventual visualization/detection (31,32). Clinical tumor samples are subjected to mild protease digestion to disrupt histones and DNA crosslinking to allow the probe access to the gene of interest. Probe is applied to the digested samples, allowed to bind or "anneal" to the genetic sequence, and the gene copy number is simply quantitated by counting the number of fluorescent signals in each cell. Many FISH systems also include a "control" probe to a centromeric gene sequence to correct for chromosomal gain or loss and to differentiate ploidy from true amplification. When this approach is used, FISH status is expressed as the ratio of the number of copies of the gene of interest to the number of control genes. In a normal situation there would be two target genes and two control genes for a ratio of 1. Because of technical issues, ratios between 1 and 2 are typically consider a "gray zone" result, with ratios ≥ 2 clearly indicating gene amplification. FISH represents a huge technical advance over older methods of gene quantification, such as Southern or DNA slotblotting, in that FISH assays can be performed on routinely processed, FFPE clinical material as opposed to fresh or frozen clinical material. In addition, the resilience of DNA makes this assay far more robust than IHC and avoids many of the problems of specimen processing and age effects that hamper protein detection assays such as IHC.

This technical advantage enabled the retrospective evaluation of FISH status as a predictor of clinical benefit in three large, pivotal Herceptin clinical trials; H0648g, H0649g, and H0650g. A total of 799 patients were enrolled in these trials, all patients tested positive at the 2+ or 3+ level using

,	in Patients Treated with	1		
		Evaluable patients	<i>Objective</i>	responses
Study	HER2 amplification	(No.)	No.	%
Study H0649g	FISH-positive	173	33	19
	FISH-negative	36	0	0
Study H0650g	FISH-positive	79	27	34
	FISH-negative	29	2	7

Table 6
Objective Response Rates by HER2 Amplification Status
in Patients Treated with Trastuzumab Alone

Table 7
Objective Response Rates by HER2 Amplification Status
for Patients Treated with Trastuzumab
and Chemotherapy vs Chemotherapy Alone—Study H0648g

	Trastuzumab and chemotherapy			Chemotherapy alone			
	Evaluable patients	Objective responses		Evaluable patients	Objective responses		
HER2 amplification	No.	No.	%	No.	No.	%	р
FISH-positive FISH-negative	176 50	95 19	54 38	168 57	51 22	30 39	<.0001 NS

the CTA. Archival tissue sections were available from 784 (98%) and FISH results were generated from 765 (96%). This high rate of technical success confirmed the robust characteristics of FISH and served to minimize some of the risk associated with retrospective analyses where missing data points can hamper the generalization of results. The first interesting finding from this analysis was that only 78% of the tumors were found to be amplified, suggesting significant discordance between an IHC score of 2+ or 3+ and the presence of HER2 gene amplification. The distribution of FISH scores shown in Fig. 2 demonstrates a bimodal distribution, with a median ratio of 6.1 in the FISH positive group and 1.2 in the FISH-negative group. This distribution is similar to other reported data and supports the cutpoint of 2 in separating amplified and nonamplified tumors (33,34).

The next series of tables and K–M graphs outline the clinical activity of Herceptin in these three trials, in both the FISH-positive and FISH-negative subgroups. Table 6 shows the single-agent response rates in studies H0649g and H0650g. Of the 65 FISH-negative patients in these two studies, objective responses were seen in only two patients and, on further testing, one

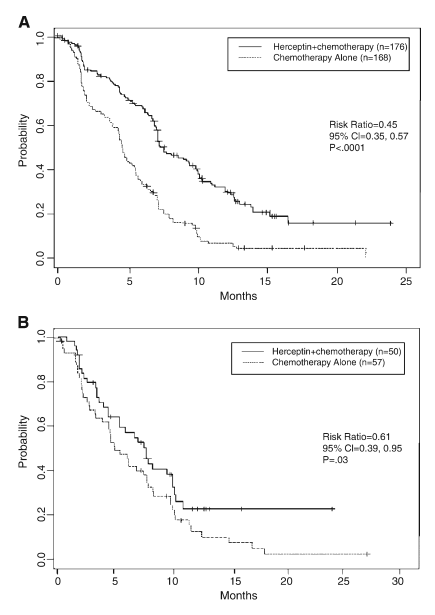


Fig. 3. Time to disease progression by HER2 amplification status for patients receiving chemotherapy + trastuzumab vs chemotherapy alone (study H0648g). (A) FISH-positive. (B) FISH-negative.

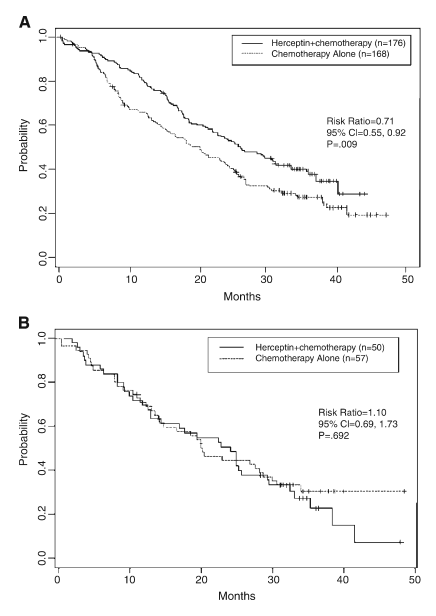


Fig. 4. Overall survival time by HER2 amplification status for patients receiving chemotherapy + trastuzumab vs chemotherapy alone (study H0648g).(A) FISH-positive. (B) FISH-negative.

of these two patients was found to have gene amplification. The same association was seen in study H0648g (*see* Table 7). In the FISH-negative group, response went from 39% to 38% with the addition of Herceptin while in the FISH-positive group response rate went from 30% to 54%.

The time to disease progression in study H0648g was strikingly improved in the FISH-positive group with a risk ratio of 0.45 (p < 0.0001); however, there was also borderline improvement for the FISH-negative group with a risk ratio of 0.61 (p = 0.03) (Fig. 3).

Finally, in evaluating the critical end point of survival there was clearly significant improvement from the addition of Herceptin in the FISH-positive population (risk ratio = 0.71, p = 0.009) that was not apparent in the FISH-negative group (risk ratio = 1.10, p = 0.692) (Fig. 4).

All the efficacy analysis in these three trials suggested that the treatment effect from Herceptin was limited to the FISH-positive population with the exception of the time to progression analysis in study H0648g. It is not clear whether this is a "real" treatment effect or whether this is a statistical anomaly, an inherent risk in the conduct of multiple retrospective subset analyses. The preponderance of the data from the other efficacy analyses, however, suggests the latter as the most reasonable conclusion.

4. CONCLUSIONS

Herceptin represents one of a few but growing class of "targeted" anticancer agents: agents that are specifically designed to precisely antagonize, or suppress, the abnormal biologic phenotype responsible for the malignant characteristics of a particular tumor. In some malignant conditions, the target biologic abnormality occurs with high frequency within the affected population, rendering screening prior to the therapeutic application of targeted therapy unnecessary. This situation is best exemplified by the abnormal *bcr-abl* tyrosine kinase in chronic myelogenous leukemia and the targeted therapeutic agent imatinib (Gleevec®). Unfortunately, this situation of "high prevalence" appears to be uncommon, with most solid tumors demonstrating significant phenotypic and/or genotypic heterogeneity. As new agents are developed targeting molecular abnormalities that affect a minority of patients with a specific tumor type, detection of the underlying abnormality and the specific methodology to detect that target will assume increasing importance when assessing therapeutic benefit. Important work to date suggests that the use of FISH to detect the primary biologic alteration, amplification of HER2, is the preferred methodology to select breast cancer patients for Herceptin therapy as compared to IHC. Diagnostic methods are inevitably destined to evolve along with remarkable advances in technology, however. This observation reinforces the vital need to prospectively incorporate tissue acquisition and archiving into ongoing clinical development plans for the new generation of targeted anticancer agents.

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10

The Epidermal Growth Factor Receptor Pathway as a Selective Molecular-Targeted Treatment in Human Breast Cancer

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CONTENTS

INTRODUCTION EXPRESSION OF THE EGF FAMILY IN BREAST CANCER EXPRESSION OF THE EGFR FAMILY IN BREAST CANCER INTRACELLULAR SIGNALING THROUGH THE EGFR FAMILY IN BREAST CANCER DEVELOPMENT AND PROGRESSION AND ROLE AS POTENTIAL MOLECULAR TARGETS FOR THERAPY REFERENCES

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SUMMARY

The regulation of normal breast development is dependent on several hormones. Among these hormones, estrogens play an essential role in the control of normal mammary development and in the etiology and progression of breast cancer.

Key Words: Breast cancer; EGF; EGFR.

1. INTRODUCTION

The regulation of normal breast development is dependent on several hormones. Among these hormones, estrogens play an essential role in the control of normal mammary development and in the etiology and progression of breast cancer. More recently, it has been recognized that normal and malignant mammary epithelial are also able to synthesize a number of different locally acting peptide growth factors that function through autocrine, juxtacrine, and paracrine pathways. Several studies have demonstrated that estrogens influence mammary epithelial cell growth both directly and indirectly by modulating growth factor production and growth factor receptor expression (1). Among these, the epidermal growth factor (EGF) family of peptides, in combination with their specific cognate receptors, is involved in the regulation of mammary gland development, morphogenesis, and lactation, and also play a pivotal role in the pathogeneses of human breast cancer (2). The purpose of this chapter is to describe the role of EGF-related growth factors and their receptors in the control of proliferation and differentiation of human mammary epithelial cells and in the pathogenesis of human breast cancer and to discuss the possibility of using drugs that selectively block the activation of the EGFR as anticancer therapy in human breast cancer.

2. EXPRESSION OF THE EGF FAMILY IN BREAST CANCER

EGF is a 6-kDa polypeptide of 53 amino acids. The human EGF gene is located on chromosome 4(3). It was originally isolated from the male mouse submaxillary gland as a factor that caused eyelid opening (4) and later from human urine as urogastrone (5). EGF is a transmembrane glycosylated protein that is biologically active as it can bind to and activate EGF receptors in adjacent cell through a juxtacrine mode of action (6). It has been found to stimulate the growth of both normal and transformed human mammary epithelial cells. Breast cancer cells are also able to synthesize EGF. In fact, EGF mRNA has been detected in a majority of human breast cancer cell lines, with the estrogen-receptor (ER)-positive breast cancer cell lines T47-D and ZR-75-1 showing higher levels of expression as compared to the ER-negative breast cancer cell lines (7). Progestins were found to specifically increase the levels of EGF mRNA in T47-D cells, while 17β-estradiol had no effect on EGF mRNA levels in this cell line (8,9). EGF mRNA was also detected in 83% of human breast tumors and in addition EGF protein has been detected in 15–30% of human primary invasive breast carcinomas (8,9). EGF is a crucial regulator of growth and differentiation in the mouse mammary gland, especially during pregnancy and lactation and during the spontaneous formation of mammary tumors (8,9).

Transforming growth factor- α (TGF- α) was first identified in the culture media of virus-transformed cells and of human tumors cell. It shares 42% homology with human EGF. Its tertiary structure is identical to EGF and is able to bind to the EGF receptor with the same affinity. Several clinical and experimental studies have demonstrated that TGF- α is an important modulator of the malignant progression of mammary epithelial cells in breast cancer (8,9). In particular, TGF- α is a potent mitogen for normal and malignant mammary epithelial cell in vitro and, similar to EGF, is able to stimulate lobuloalveolar development of the mouse mammary gland in vivo (8,9). The basal levels of TGF- α are higher in ER-positive estrogen-responsive, breast cancer cell lines. TGF-a mRNA and protein are induced by physiological concentration of 17B-estradiol. The mechanism of induction seems to be transcriptional because several imperfect estrogen response elements (EREs) that are located in the promoter region of the TGF- α gene are thought to be active, as determined by an increase in CAT activity of transiently transfected breast cancer cells. This induction can be blocked with the antiestrogen tamoxifen, suggesting that this effect is mediated through the ER. In this regard, it has been demonstrated that tamoxifen can reduce the production of TGF- α by 30–70% in primary human breast tumors that are ER positive (8,9). In addition, in one study tamoxifen treatment of breast cancer patients resulted in a 10-fold reduction in TGF- α tumor levels. These data demonstrate that TGF- α can act as an autocrine growth factor and can function as a mediator in part for the growthpromoting effects of oestrogen in human breast cancer cells. The majority of the tumors that express higher levels of TGF- α also express higher levels of the EGF receptor (EGFR), suggesting that an autocrine loop may be operative. TGF- α mRNA and protein levels have been detected in 40-70% of human breast tumors. RNA in situ hybridization has shown that TGF- α mRNA is expressed mainly in breast cancer cells and not in the surrounding stromal cells or in infiltrating lymphoid cells (8,9). TGF- α has also been detected in premalignant atypical ductal hyperplasias, ductal hyperplasias, ductal carcinoma *in situ*, and in 30–50% of primary and metastatic human breast carcinomas at levels that are generally two to threefold higher than the levels found in benign breast lesions or in normal mammary tissues (8,9). TGF- α has also been found in the pleural effusions and in the urine samples from metastatic breast cancer patients. Higher levels of TGF- α in pleural effusions correlate with poor prognosis and performance status and with tumor burden. TGF- α mRNA has been detected during the proliferative, lobular development of both the rat and human mammary gland and its expression is enhanced during pregnancy and lactation, suggesting some form of hormonal control of TGF- α production in vivo (8,9).

Amphiregulin (AR) is a glycoprotein that is structurally related to EGF, but it has lower affinity for the EGFR than TGF- α and EGF. Expression of high levels of endogenous AR mRNA and protein has also been found in several normal human mammary epithelial cell strains in culture and in nontransformed human mammary epithelial cell lines. Furthermore, AR and TGF- α can function as autocrine growth factor in these cells. AR has a bifunctional mode of action. Depending on the concentration, the presence of other growth factors and the target cells, it can either stimulate or inhibit cell proliferation. Although AR has been reported to inhibit the growth of some breast cancer cell lines, there is also evidence to suggest that AR can function as autocrine growth factor in human breast cancer cells. In fact, AR is expressed in a number of human ER-positive and ER-negative breast cancer cell lines (8,9). ER-positive breast cancer cell lines generally express higher levels of AR when compared to ER-negative cell lines, and estrogen treatment induces the expression of AR mRNA in MCF-7 breast cancer cells. A significant correlation between AR and ER expression was found in a subset of human primary breast carcinomas that were examined for AR mRNA expression by Northern blot analysis. AR mRNA expression was found in approx 60% of these tumors, and all the tumors that were positive for AR expression were also found to be ER positive.

The possible role of heregulins (HRGs) in regulating the proliferation and the differentiation of human breast cancer cells has not yet been clarified, as HRGs have been demonstrated to either stimulate or inhibit the growth breast cancer cells in vitro and to induce the differentiation of breast cancer cells at least with respect to facilitating the expression of milk protein. HRG mRNA has been detected in both normal and malignant breast tissues and in a small number of human breast cancer cell lines (8,9). Its distribution is more restricted than that of TGF- α or AR, because generally ER-negative, estrogen-independent cell line express HRG, whereas only one ER-negative breast cancer cell line was found to express low levels of HGR mRNA (8,9). However, HRG seems to inhibit the effect of estrogen in ER-positive estrogen-responsive breast cancer cells (8,9). It has been show that HRG can stimulate lobular–alveolar development and the production of milk proteins in the mouse mammary gland and that antisense oligonucleotides against HRG can abolish branching morphogenesis and lobular alveolar differentiation of the mammary gland (8,9).

3. EXPRESSION OF THE EGFR FAMILY IN BREAST CANCER

The EGFR family consists of four closely related genes: c-erb-B1 or EGFR, c-erb-B2 or HER2, c-erb-B3 or HER3, and c-erb-B4 or HER4. These receptor proteins are glycosylated and share a similar primary structure consisting of an extracellular, ligand-binding domain (which has two cysteine-rich regions), a single transmembrane region, a short juxtamembrane sequence, and an intracellular domain that contains a tyrosinekinase domain flanked by a large hydrophilic carboxyl tail. The carboxyl tail displays sequence heterogeneity and carries several tyrosine autophosphorylation sites. The four members of the EGFR family have a high degree of homology in the tyrosine kinase domain, while the extracellular domains are less conserved, which is indicative of different specificity in ligand binding. Different classes of ligands that bind to and activate distinct sets of individual receptors can be distinguished within the EGF-like growth factors. The first group, that includes EGF, TGF- α , and AR, can exclusively bind to EGFR. Heparin binding EGF (HB-EGF) and betacellulin (BTC) can efficiently interact with both the EGFR and c-erb-B4, whereas epiregulin is a broad-spectrum receptor ligand. Finally, HRGs can bind to erb-B3 and erb-B4. Following ligand binding, these receptors dimerize, which may result in either homodimerization or heterodimerization. Dimerization results in tyrosine kinase autophosphorylation, which then activates a number of different intracellular signaling pathways (10,11). The signaling pathway involves activation of ras and mitogen-activated protein kinase (MAPK), which activates several nuclear proteins required for cell cycle progression from the G₁ to the S phase. This activation is critical not only for cell proliferation. Several studies have demonstrated that EGFR-mediated signals also contribute to other processes that are crucial to cancer progression, including angiogenesis, invasion, metastasis, and inhibition of apoptosis (12). In cells that express EGFR and c-erb-B2, any of EGF agonists will induce formation of EGFR/c-erb-B2 heterodimers, as well as EGFR/EGFR homodimers. This cross-activation extends to most of the receptor combinations, so that activation of one receptor will generally lead to some activation of other coexpressed EGF family receptors (13, 14). Heteromerization with other *erbB* family receptors is required for the activation of c-erb-B3, which is devoid of intrinsic catalytic activity (15). c-erb-B2 is an orphan receptor, because none of the known ligands bind to c-erb-B2. c-erb-B2 is activated as heterodimers through interactions with

other EGF family receptors following ligand binding and as homodimers in cancer cells with an amplified and overexpressed c-*erb*-B2 gene (16).

The EGFR family is involved in the regulation of mammary growth and differentiation. The female mammary gland undergoes extensive postnatal development under the influence of systemic hormones, including EGF family ligands and their cognate receptors. All four erb-B family receptors are expressed in mammary glands of adult females, but the EGFR and c-erb-B2 are preferentially expressed in young females (17–19). The first postnatal episode of mammary development occurs at puberty, and leads to elongation and branching of the mammary ducts to extend throughout the fatty mesenchyme. EGFR and c-erb-B2 are present both in the stroma and the epithelium and are tyrosine phosphorylated, which is indicative of signaling activity (18,19). The EGFR is important at puberty, because expression of dominant-negative EGFR impairs ductal morphogenesis (20). The second wave of activation of *erbB* family receptors occurs in pregnancy (18,19). Dominant-negative c-erb-B2 and c-erb-B4 transgenes interfere with lobuloalveolar expansion and milk protein production early and late in the postpartum period, respectively, which is consistent with the ability of neuregulin-activated c-erb-B2 to drive mammary differentiation (17,21-23). All four receptors have been reported to be expressed to some extent also in breast tumors. EGFR expression is generally found between 30% and 50%, although it can range from 14% to 91%, depending on the method of assessment (24,25). EGFR positivity is also common in ductal carcinoma in situ (DCIS) (26). In breast tumors, EGFR overexpression is almost never caused by gene amplification; rather, it is a result of increased receptor synthesis (27). Overexpression of c-erb-B2 is reported in approx 30% of breast tumors (28) and it is generally associated with gene amplification (29,30). Amplification and/or overexpression of c-erb-B2 occurs more frequently (in up to 60% of cases) in DCIS (31,32). c-erb-B3 is overexpressed in approx 20% of infiltrating breast cancers. As gene amplification has not been observed, overexpresion is most likely to be a result of increased transcription (33). Overexpression of c-erb-B4 is relatively uncommon in breast cancer, and *erb*-B4 expression may be suppressed in carcinoma (34–37).

4. INTRACELLULAR SIGNALING THROUGH THE EGFR FAMILY IN BREAST CANCER DEVELOPMENT AND PROGRESSION AND ROLE AS POTENTIAL MOLECULAR TARGETS FOR THERAPY

Because activation of the TGF- α /EGFR autocrine growth pathway is a common mechanism for autonomous, dysregulated cancer cell growth in the most common types of human epithelial cancers, the EGFR-driven

autocrine pathway is a rational target for cancer therapy. The design of anticancer therapies using a selective EGFR inhibitor was proposed by Mendelshon in the early 1980s as one the first approaches for interfering with a specific cancer cell molecular target (38). In the past 20 yr, a large body of experimental and clinical studies have supported this hypothesis. In this respect, the identification of selective and potent inhibitors of EGFR and of c-erb-B2 that can be developed as anticancer drugs has been one of the most successful examples of translation research in cancer. Trastuzumab (Herceptin®), a humanized monoclonal antibody generated against the c-erb-B2 protein, has shown activity in c-erb-B2 overexpressing metastatic breast cancer (39,40), and enhances survival when given in combination with chemotherapy (41). Two anti-EGFR therapeutic approaches have been shown potentially effective in clinical trials: monoclonal antibodies (MAbs) and small molecule inhibitors of the EGFR tyrosine kinase enzymatic activity. MAbs are raised against the extracellular domain of the EGFR to block ligand binding and receptor activation. Tyrosine kinase inhibitors (TKIs) prevent the autophosphorylation of the EGFR intracellular tyrosine kinase domain. The molecules are generally reversible competitors of ATP for binding to the intracellular catalytic domain of the EGFR tyrosine kinase. The most promising small molecule selective EGFR-TKIs belong to three series of compounds: 4-anilinoquinazolines, 4-ar(alk)ylaminopyridopyrimidines, and 4-phenylaminopyrrolopyrimidines. Both anti-EGFR MAbs and EGFR-TKIs have shown efficacy in relevant preclinical models, such as human cancer cell lines in vitro and in vivo. The blockade of EGFR signaling in cancer cells determines not only inhibition of cell proliferation but also other effects by inhibition of cell proliferation that could be relevant in the clinical setting, such as antiangiogenetic effects by inhibition of tumor cell production of proangiogenetic growth factors and possibly by direct cytotoxicity on endothelial cells in tumor vessels such as antiinvasive and antimetastatic effects.

DCIS is a preinvasive breast lesion that accounts for 30% of screeningdetected breast cancer (42). Untreated DCIS progresses to invasive breast cancer in 25–30% of patients (43–45). DCIS express both EGFR and c-*erb*-B2 which might play an important role in proliferation and progression of DCIS. Chan et al. (46) have investigated the effects of an anti-c-*erb*-B2 monoclonal antibody (trastuzmab, Herceptin) and of an anti-EGFR selective TKI (Gefitinib, Iressa) on in vivo growth of ER-negative DCIS. These authors have detected no significant effects on epithelial proliferation by trastuzmab treatment, while gefitinib decreases proliferation and activates apoptosis in DCIS. These data show that gefitinib may have potential as an adjuvant therapy for the treatment and chemoprevention of DCIS.

EGFR and c-*erb*-B2 overexpression has been also associated with cancer cell resistance to antineoplastic agents (47–51). In this respect, the transfec-

tion of human breast cancer cells with an EGFR expression vector results in the development of resistance to several chemotherapeutic agents (52). Moreover, human breast cancer cell lines that become resistant to various cytotoxic agents overexpress EGFR, c-erb-B2 and various ligands, including TGF- α (53). Signaling from EGFR, via the mitogen-activated protein kinase (MAPK) pathway, is believed to protect breast cancer cells against ionizing radiation and to be involved in rapid repopulation following radiotherapy (47,48,54). Consequently, disrupting EGFR signaling with selective anti-EGFR drugs should result in cellular radiosensitization and enhanced sensitivity to chemotherapeutic agents. In addition, it could slow repopulation after radiotherapy or chemotherapy (54). It is conceivable that the cellular damage induced by chemotherapy or by ionizing radiation can convert EGFR ligands from growth factors into survival factors for cancer cells that express functional EGFR. In this situation, the blockade of EGFR signaling in combination with cytotoxic drugs or with radiotherapy could cause irreparable cancer cell damage leading to increased programmed cell death. The enhancement of anticancer activity of conventional cytotoxic treatments, by interfering with EGFR activation, may have relevant clinical implications. In this respect, treatment with conventional doses of cytotoxic drugs or of radiotherapy in combination with signal transduction inhibitors, such as anti-EGFR-selective agents, could be an effective novel anticancer strategy, which is less toxic and more tolerable than other clinical approaches for increasing the activity of cytotoxic drugs or radiotherapy (55,56). Clinical trials are ongoing in a variety of tumor types, although the results of some of these trials have been controversial. However, the combined or the sequential use of EGFR-targeted agents and chemotherapy is currently under clinical investigation in human breast cancer. The effect of combining EGFR-targeted agents with radiotherapy has not been examined in detail in breast cancer, although it has been shown to be effective in other types of cancer (57). A recent study, however, has reported that breast cancer cells expressing a dominant-negative EGFR were more sensitive to radiation, suggesting that EGFR disruption could increase the sensitivity to radiation therapy (54). EGFR expression in breast cancer is generally associated with an increased likelihood of failure to respond to endocrine therapy (58–61). A similar relationship between c-erb-B2 overexpression, a poor prognosis, and hormonotherapy resistance has also been observed, although these associations as yet remain controversial (58,60-63). The concept that peptide growth factors can act as the growth mediating factors in the growth of hormone-sensitive breast cancer is not a new one. It had its origins in the late 1980s, when it was first recognized that estrogens were able to stimulate expression of a number of growth factors, including TGF- α , in hormonesensitive human breast cancer cell lines (64,65). Numerous studies have shown that key receptors in such pathways (e.g., ERs) are subject to activation by both estrogens and peptide growth factors (66–68). The important significance of such convergence in hormone-sensitive breast cancer cells is that antihormonal drugs not only possess antiestrogenic activity through their ability to block ER signaling, but also have antigrowth factor actions by virtue of their ability to disrupt the intimate cross-talk between estrogen and growth factor signaling (69,70). Expression of EGFR and c-erb-B2 is suppressed by long-term therapy with estrogens in vitro (71-74). These data suggest that hormone-sensitive breast cancer cells possess control mechanisms to limit EGFR/erb-B2-mediated signaling. This concept may have significant clinical implications. It has been observed a time-dependent increases in the expression of EGFR and c-erb-B2 after antiestrogen treatment of MCF-7 human breast cancer cells in vitro. MCF-7 cells are estrogen responsive for their growth and are growth inhibited by many antiestrogenic drugs. However, their continuous culture in the presence of tamoxifen or faslodex eventually generates sublines that tolerate the presence of the antiestrogens, regrowing at rates equivalent to the original hormoneresponsive parenteral cells (75). McClelland et al. (76) have shown that antihormonal-resistant MCF-7 sublines express up to 10-fold increased amounts of EGFR mRNA and protein. A parallel increase in c-erb-B2 immunostaining in the antiestrogen-resistant MCF-7 cells was also observed. Furthermore, treatment with gefitinib of tamoxifen- or faslodexresistant MCF-7 breast cancer cells determines inhibition of EGFR autophosphorylation and of cell growth, indicating that the autocrine EGFR loop is critical to the growth of these antihormonal-resistant cells. The increase in cellular expression of EGFR in hormone-resistant breast cancer cells thus appears to provide a promising molecular target for the effective treatment of endocrine-resistant human breast cancer. McClelland et al. have also shown that combination therapies with simultaneously target estrogen (tamoxifen) and EGFR signaling (gefitinib) may be more effective than the sequential use of such drugs. Moreover, these experimental studies suggest that anti-EGFR treatment of hormone-responsive breast cancer may contribute to prevention of the development of the hormoneresistant tumors.

There is experimental evidence that EGFR and c-*erb*-B2 coexpression confers a more aggressive clinical behavior (77–79). The rationale for targeting simultaneously the two receptors stems from their frequent coexpression in breast cancer and their capacity to form heterodimers that activate signal transduction pathways. Four independent research groups have provided elegant experimental data showing that the EGFR-selective tyrosine kinase inhibitor gefitinib has potent in vitro and in vivo antitumor activity against human breast cancer cell lines that express the EGFR and that overexpress c-*erb*-B2 (80–83). Gefitinib treatment efficiently blocks EGFR autophosphorylation and c-*erb*-B2 transphosphorylation (80–83).

This effect is followed by inhibition of both MAPK and PI3K-akt signaling in breast cancer cells with subsequent growth inhibition and induction of apoptosis (81,82). Furthermore, the combined treatment with gefitinib and trastuzumab is frankly synergistic both in inducing cell growth inhibition and apoptosis in vitro and in vivo (81, 82), suggesting that the simultaneous targeting of the EGFR and c-erb-B2 may result in an efficient tumor growth inhibition in breast cancer patients whose tumors coexpress both receptors. A possible biochemical and pharmacological explanation for the activity of gefitinib in EGFR and c-erb-B2 overexpressing breast cancer cells has been recently provided by Anido et al. (83), who have demonstrated that gefitinib treatment blocks EGFR activation and also induces the formation of inactive EGFR/c-erb-B2 and EGFR/c-erb-B3 heterodimers and prevents HRG signaling. In this situation, gefitinib treatment significantly blocks signaling through the entire EGFR family of receptors because it also reduces the levels of c-erb-B2/c-erb-B3 that are potentially activable by HRG (83,84). Collectively, these experimental evidence provide a strong rationale for studying the combination of gefitinib and trastuzumab in breast cancer patients who coexpress both the EGFR and c-erb-B2.

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11

p53 as a Prognostic and Predictive Indicator

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CONTENTS

INTRODUCTION: WHAT WENT WRONG WITH *p53*? METHODOLOGICAL VARIABILITY ARE ALL GENE MUTATIONS OF CLINICAL RELEVANCE? *p53* AND RESPONSE PREDICTION *p53* AND PROGNOSIS CONCLUSIONS REFERENCES

SUMMARY

Cancer is a disease of genes, and the p53 gene appears to be the most frequently mutated gene associated with human cancer. p53 is crucially involved in numerous central pathways determining the life and death of a cell. The information about the functional status of p53 has been considered to be extremely important for estimation of prognosis and the need of further treatment as well as for prediction of response and the selection of the optimal therapy. The aim of this chapter is to evaluate why one of the most promising and most intensively studied markers is not recommended for routine clinical use to date. The pitfalls of the p53 analysis and the challenges of well-designed p53 studies are elucidated and the need of carefully planned studies using an optimal p53 analysis method is pointed out.

Cancer Drug Discovery and Development: Biomarkers in Breast Cancer: Molecular Diagnostics for Predicting and Monitoring Therapeutic Effect Edited by: G. Gasparini and D. F. Hayes © Humana Press Inc., Totowa, NJ **Key Words:** *p53*; biological marker; prognostic; predictive; *p53* sequencing; *p53* immunohistochemistry; *p53* study design.

1. INTRODUCTION: WHAT WENT WRONG WITH *p53*?

The treatment of breast cancer, a genetically heterogeneous malignant disease, has been greatly improved by providing individual treatment for the patient. While prognostic factors determine the need for further treatment, predictive factors are needed to select the appropriate therapy with the greatest likelihood of eliciting a response.

From a biological aspect, p53 should a powerful predictive and prognostic marker. A p53 has been shown to be responsible for the activation of the apoptotic machinery leading to cell death. The effect of most standard chemotherapies, tamoxifen, and also radiation treatment, is based on the induction of p53-dependent apoptosis. Thus, a defect in the p53 pathway would cause resistance to treatment. Predicting either resistance or response to treatment is a key feature of a predictive marker. Furthermore, p53 potently inhibits cell growth and provides protection from malignant progression. Taken together, these data suggest that p53 should represent an independent prognostic marker.

However, in a review of *Molecular Markers in the Treatment of Breast Cancer* published in 2000, Hamilton and Piccart concluded that currently available data do not support the use of p53 as a predictive factor in the therapy of breast cancer (1). The prognostic impact of p53 has been reported in numerous publications, which have been summarized in the 1999 consensus statement of the College of American Pathologists. Here, p53 has been ranked as a category II factor, defined as "having been extensively studied but whose importance remains to be validated in statistically robust studies" (2).

So what went wrong with *p53*?

2. METHODOLOGICAL VARIABILITY

2.1. p53 Immunohistochemistry (IHC)

2.1.1. P53 IHC—BACKGROUND

The p53 protein is normally present at extremely low levels. One action of p53 is suppression of cellular proliferation; rapid degradation of the p53 protein after synthesis, regulated to a large extent by Mdm2, allows normal cells to grow (3,4).

Stabilization of the p53 protein occurs as a common response to cellular stress including DNA damage, hypoxia, oncogene activation, pH, and temperature changes (heat shock). In response to these signals, p53 becomes active as a result of stabilization and mediates the inhibition of cell growth

via cell cycle arrest, DNA repair, or cell death, to prevent the development and progression of malignant cells.

2.1.2. SIGNIFICANCE OF IHC RESULTS

Wild-type p53 protein has a very short cellular half-life. In contrast, mutated p53 appears relatively resistant to Mdm2-mediated degradation. Therefore, wild-type p53 protein is rarely detectable by routine IHC staining.

Mutation of the p53 gene can result in the failure of normal degradation of p53 (3). These proteins are dysfunctional, in contrast to proteins that are physiologically stabilized as a response to cellular stress. Because IHC detects stabilized p53 protein without further specification, this method is fraught with false-positive results, as it does not differentiate between pathological (due to gene mutation) and physiological (due to cellular stress) overexpression.

The majority of publications over the last 15 yr report IHC results founded on the assumption that the overexpression of p53 in tumors occurs due to gene mutation. The frequency of p53 overexpression in breast cancer ranges from 40% to 57% (5), which is much higher than the frequency of *TP53* gene mutations in breast cancer (20–30%).

Comparative studies of IHC vs gene sequencing show that the concordance between the two methods is only 12-30% (6–9).

IHC may also produce false-negative results. For instance, changes in the protein structure affecting the antibody-binding sites due to gene mutation; or the formation of a premature STOP codon most frequently occurring as a consequence of deletion or insertion mutations, causing a shift of the reading frame and preventing gene transcription and translation into protein at all.

Moreover mutations that do not result in greater expression of the p53 protein are termed "null mutations" and have been reported to occur at a frequency of 31% in breast cancer (10). These may or may not alter p53 function.

2.1.3. STANDARDIZATION OF IHC

A number of antibodies to detect different epitopes of the p53 protein are available. Most groups use Mab 1801 (Oncogene Science) or DO-7 (DAKO), or both, for p53 IHC. The staining protocols, detection, and reporting procedure for positive staining are not standardized. A number of parameters are variously reported: the staining intensity, the percentage of tumor cell nuclei, the staining index (=product of staining intensity and area), or the staining score (=sum of staining intensity and positive tumor cell nuclei). Different cutoff points ranging from 0% to 75% further complicate the issue (11,12). The role of interobserver variability, which is linked to microscopic detection methods, can be estimated only after the method itself has been standardized.

Although p53 IHC promises to be a simple and inexpensive method, it currently bears all the major disadvantages of a marker analysis method with respect to significance and standardization. This is a major hindrance to correct estimation of the clinical importance and the independence of p53 IHC as a marker. In the absence of comparative studies, it is currently not even clear whether the standardization of IHC can improve the situation.

2.2. Direct Detection of p53 Genetic Abnormalities

2.2.1. Screening Methods

Various screening methods have been applied in different studies, promising high-throughput screening for mutations. These are based on the analysis of sequence-dependent changes in the conformation of single- and double-stranded DNA using capillary electrophoresis (13,14). The specificity, sensitivity, and reproducibility of these methods are controversial. In some studies, when compared to direct sequencing, these methods will not detect up to 50% of mutations. Moreover, these methods are unable to distinguish between polymorphisms and relevant mutations (15,16). In most articles, screening results are compared with the clinical outcome. Consecutive sequencing is performed only for tumors which proved to be positive at initial screening. Tumors with negative screening are disclosed from sequencing and often incorrectly adjudged to have a normal genotype. In this regard, the rate of detected p53 abnormality using the most common of these methods, SSCP, is 20%, but the concordance of SSCP with sequencing is only 50%. Thus the mutation frequencies derived from studies using screening methods prior to sequencing may not reflect true frequencies, as a significant fraction of mutations may have been missed.

Commercially available array-based technologies are gaining popularity as a screening method for mutations (17, 18). However, comparative studies concerning sensitivity, specificity, and reproducibility have not yet been published. To promote this technology, one has to carefully design studies that permit comparisons with the most sensitive technique at the present time, namely p53 sequencing.

2.3. p53 Sequencing

The p53 tumor suppressor gene encodes a nuclear phosphoprotein that directly binds to DNA. Directly interacting with DNA as a transcription factor, p53 controls numerous genes involved in DNA repair, cell division, and apoptosis. To allow abnormal proliferation and progression, cells dedicated to become malignant have to be deprived of their tumor suppressor ability. The most commonly used pathway to inactivate p53 in human cancer is mutation of the gene itself. In breast cancer 20% of tumors appear to be mutated.

2.3.1. STANDARDIZATION OF SEQUENCING

Although IHC or SSCP are more appealing methods to evaluate p53 abnormalities, the preceding paragraphs illustrate the inaccuracies of these approaches. Currently, the gold standard to detect p53 abnormalities is sequencing of the gene.

Sequencing has the reputation of being time consuming and cost ineffective. Today, however automated capillary sequencing provides a high throughput, a computerized standardized interpretation, and characterization of mutations (6-8).

Sequencing results can either be positive (mutated) or negative (normal, wild-type). Thus, problems related to different scoring systems, cutoff points, and interobserver variability do not exist. For p53 analysis it is now generally accepted that, to achieve highest sensitivity, direct sequencing has to be applied to the entire p53 gene including splice sites. In older studies, sequencing was limited to hot spot regions spanning exons 5–8. These studies might not reflect true frequencies, in consideration of the fact that up to 20% of mutations are missed in breast cancer (24).

Reproducibility and correct interpretation are ensured by a standardized number of reanalyses and by sequencing the opposite DNA strand in order to confirm mutations. Routine coamplification of positive and negative controls has disclosed problems related to the contamination or fidelity of the DNA polymerase (enzyme) used for polymerase chain reaction. Characterization of mutation ensures standardized reporting of mutations and identification of polymorphisms.

In contrast to IHC, sequencing is highly sensitive and specific and can be applied to various materials including fine-needle aspirates and paraffinembedded and frozen tissue (25).

Sequencing will not detect inactivation of p53 by mechanisms other than mutation, for example, cytoplasmatic sequestration: p53 protein imprisoned in the cytoplasm fails to activate transcription of target genes and cannot mediate cell cycle arrest or apoptosis. To date there are no studies demonstrating the frequency and cancer-specific relevance of such finding. It is unclear whether the sequestration is stable and what happens when p53is up-regulated in response to DNA damage. So far mutation of the p53 gene, and not cytoplasmatic sequestration of the protein, is the most commonly known path to inactivate the p53 tumor suppressor function in human cancer.

3. ARE ALL GENE MUTATIONS OF CLINICAL RELEVANCE?

3.1. Significance of p53 Gene Mutation

In multivariate analysis, the presence of *TP53* mutations is consistently associated with an increased risk of relapse and death from breast cancer

(19). In addition, a mutant p53 genotype appeared to be clearly associated with a lack of response to DNA-damaging treatment in different cancer types, including breast cancer (6–8,10) (Tables 1; [8,10,16,20–23] and 2 [6–8]).

However, it has become increasingly clear that not all *p53* mutations are of equal significance, almost certainly owing to the pleiotrophic features of this gene and its protein. Certain data indicate functional differences between the central and peripheral coding regions. The highly conserved region located in the middle of the protein is responsible for DNA binding and is affected most frequently by mutations (80%).

There are some reports of mutations that show that the L2 and L3 loops of the p53 gene may have a distinct tumorigenic potential (26). In contrast, only 4% of mutations occur in the tetramerization domain. However, for this region, germline mutations in Li–Fraumeni cancer syndrome families have been published, proving the clinical relevance of such rare mutations (27).

A specific sensitivity to chemotherapeutic agents has been reported for certain mutations (28). Even if the different types of mutations as well as the different location of mutations confer a specific grade of treatment sensitivity, clinically this does not seem to impair the prognostic and predictive implications of p53. While the response to (p53-dependent) treatment was shown to be related to the p53 genotype in more than 168 patients from neoadjuvant studies, we were unable to identify a single tumor that responded to treatment in the presence of a p53 mutation (Table 2; [6-8]). This was found to be true for more than 70 different mutations in the p53 gene. The results were reproducible in three different tumor types (breast, colorectal, lung cancer).

4. p53 AND RESPONSE PREDICTION

4.1. Biologic Rationale

The effect of the majority of currently used chemotherapeutic drugs is based on the induction of DNA damage, which favors rapidly growing cells. These drugs share a final common pathway: apoptosis. p53 plays a central role in this pathway: DNA damage is the strongest promoter of p53 activation. The latter is responsible for the induction of apoptosis in response to DNA damage (29).

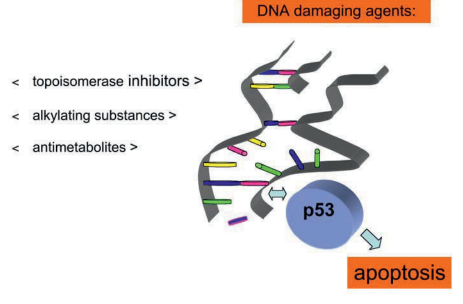
Therefore, in several preclinical studies, *TP53* gene mutation represents a crucial defect in the apoptosis pathway and results in drug resistance (30,31).

These studies suggest that *p53*-mediated drug resistance is important for several agents, including anthracylines (doxorubicin, epirubicin), antimetabolites (5-fluorouracil, methotrexate, gemcitabine), topoisomerase blockers (CPT-11), alkylating substances (cyclophosphamide, ifosfamide, cisplatin), hormonal therapies such as tamoxifen and aromatase inhibitors, and radiation therapy (*32*) (Fig. 1).

		No.		Technical				Relation of p53	
Authors (ref.)	Journal	of patients	Technique	of patients Technique specificities	Therapy	Design	Prospective	Design Prospective and response p-value	p-value
Anelli (20)	Anelli (20) Ann Oncol 2003	73	Sequencing		Paclitaxel + docorubincin	Neoadjuvant	Yes	Predictive	0.004
terns (10)	Berns (10) Cancer Res 2000	202	Sequencing	Sequencing Exons 2-11	TAM	Metastatic	No	Predictive	0.0001
Candioler (8)	Kandioler (8) Clin Cancer Res 2000	25	Sequencing	Sequencing Exons 2–11 + FEC splice sites	FEC	Neoadjuvant	No	Predictive	0.0029
(16) (16)	Elledge (16) Breast Cancer Res Treat 1998	360	IHC	Pab 1801	TAM	Metastatic	No	No	
Aakris (21)	Makris (21) Clin Cancer Res 1997	80	IHC	PAb240, PAb1801	Various therapies Neoadjuvant	Neoadjuvant	No	No	
ormenti (22)	Formenti (22) 1J Radiat Oncol Biol Phys 1997	35	IHC		5-FU + RT	Neoadjuvant	Yes	Predictive	0.01

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	Kelationship of p_{λ}	5 Sequencing	Kelationship of <i>p33</i> Sequencing and Kesponse to Neoadjuvant 1 reatment in Cancer Patients No. Converting Sequencing result	adjuvant 1	Sequenci	eatment in Cancer Sequencing result	Patients	Immunohis	Immunohistochemistry
Author (ref.)	Journal	of patients	of patients neoadjuvant therapy Response Mutant Normal p-value	Response	Mutant	Normal	p-value	Positive	Positive Negative
Kandioler et al. (8) Clin	Clin Cancer Res 2000	25	Breast cancer 5-FU/epirubicine/ cyclophosphamide	CR PR SD PD	0	$\frac{18}{10}$	0.0029	$\frac{1}{14}$	17 3
Kandioler et al. (7) JTCVS 1999	JTCVS 1999	58	Lung cancer Cisplatin/ifosfamide	CR PR SD PD	$\begin{array}{c} 0\\ 18 \end{array}$	32 8	0.001	9^a	7a 3a
Kandioler et al. (6) Ann Surg 2002	Ann Surg 2002	68	Rectal cancer Short-term radiation	CR PR SD PD	0 29	14 21	0.000	na na	na na



Doxorubicin, Cyclophosphamid, 5-FU, Methotrexate, Gemcytabine Radiation... Fig. 1. Functional *p53* is required to respond to DNA-damaging substances.

Fig. 1. Functional p53 is required to respond to DNA-damaging substances. DNA damage is the most important trigger for p53 transcription. The p53 protein binds directly to DNA and initiates apoptosis. Loss of p53 function will therefore result in treatment resistance.

4.1.1. p53-Independent Substances

Paclitaxel exhibits significant antitumor activity in breast cancer patients in prospective randomized studies (33). In vitro experiments show cells lacking functional p53 to be particularly sensitive to paclitaxel (34). Because paclitaxel blocks cells in the G₂/M phase of the cell cycle, this agent has been proposed as a radiosensitizer.

In a small analysis of a breast cancer patient cohort treated neoadjuvantly with paclitaxel monotherapy, we were able to show in the clinical setting that the response to paclitaxel was positively related to the presence of *TP53* gene mutations (detected with sequencing), and was not related to apoptosis (detected with the TUNEL assay), suggesting that cell death caused by paclitaxel occurs in a *p53*-independent manner (8). As the response to paclitaxel occurred preferentially in *p53* mutants, we hypothesized that it might be the defect in the G1 cell cycle checkpoint which increases sensitivity to taxanes in *p53* mutants, probably due to accumulation of cells in the phase of mitosis (Fig. 2).

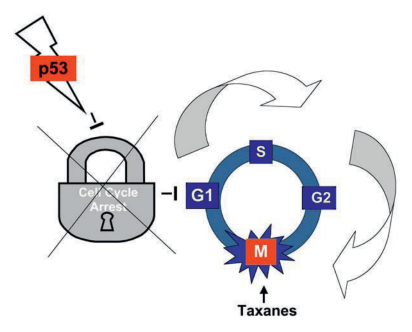


Fig. 2. Mutant, dysfunctional p53 may improve response to taxanes. During mitosis, taxanes are effective against tumor cells, inducing a p53-independent cell death. Tumor cells harboring a mutant p53 gene will not be arrested in the G₁ phase. These cells remain unobstructed and can reach the phase of mitosis in a more synchronized way. Therefore mutant tumor cells may respond even better to taxanes.

4.2. Study Design

As predictive markers are used to estimate the likelihood of sensitivity or resistance to a distinct therapy, the end point of studies evaluating predictive markers should be the response to treatment (35). The neoadjuvant setting is ideal to measure tumor response directly. Adjuvant trials offer only indirect parameters of response, namely disease-free and overall survival. In the adjuvant setting, a survival advantage can be attributed only indirectly to treatment response, on the condition that an untreated control group is coevaluated. An untreated control arm is not needed in neoadjuvant trials, as there is unlikely to be a response to no treatment. In this regard, several studies have shown that complete pathologic response after several cycles of therapy is highly associated with disease-free and overall survival (39,40). Therefore retrospective p53 analyses from patients who are prospectively entered into neoadjuvant trials might achieve a high level of evidence (see clinical utility score of tumor markers) (35–37). Such studies are cost

effective and time saving, provided all patients of the initial study can be included.

Berns reviewed 26 reports from 1995 to 2000 that correlated p53 status and response to treatment, including nearly 6000 breast cancer patients. This review included five studies using p53 sequencing analysis, but unfortunately these were all adjuvant trials and, in addition, patients were treated with multiple agents. All five sequencing studies were reported as being predictive of response to tamoxifen or p53-dependent chemotherapy. The results from the 21 reviewed IHC studies were conflicting (6 predictive, 15 not predictive). The reviewed studies included four neoadjuvant trials, all of which unfortunately used IHC for p53 analysis and were all negative for the prediction of response (10,21,43).

Furthermore, the review included five trials with second-line treatment in the metastatic setting, which also allows direct response assessment, although most frequently clinical and not pathohistological response is evaluated in such studies. However, in the two studies which used sequencing, the p53 genotype was reported to be predictive. In the three IHC studies p53 results did not provide predictive information (10).

Currently, no reported studies of p53 as a predictive factor takes into account all of these issues; monotherapy, evaluation of a reliable end point, and issues regarding the technical aspects of the assay for p53 abnormalities (35-37).

In Table 1 we reviewed recent studies that meet the minimum requirements to assess the predictive value of p53 in breast cancer. These studies provide a neoadjuvant or metastatic setting, allowing a direct response assessment.

All studies using p53 sequencing technique proved to be predictive of response to p53-dependent therapies. Results from the listed IHC studies were found to be inconsistent; the majority were not predictive for response. However, most of the studies were retrospective and did not include the complete cohort from the initial prospective trial.

In three retrospective studies on three different cancers we evaluated the predictive power of p53. Using p53 sequence analysis in neoadjuvantly treated patients only, we found that the p53 genotype consistently and completely demarcated a group of patients who did not respond to treatment (Table 2; [6-8]). Sequencing analysis of these 168 patients who had received p53-dependent neoadjuvant treatment revealed no single patient harboring a p53 mutation among those who responded to treatment. The probability that a mutant p53 genotype indicates treatment failure (=specificity) was 100% in breast, lung, and rectal cancer, respectively. In these studies the p53 genotype proved to be a powerful predictor of treatment failure. The complete identification of a group of patients (i.e., responders or nonresponders) is a prerequisite for the clinical utility of a predictive marker, as it allows the clinician to make treatment decisions (44).

5. *p53* AND PROGNOSIS

The p53 gene appears to be involved in a vast number of essential cellular functions. As a transcription factor it binds to a number of genes, either turning them on or off. Cells with inactivated p53 generally exhibit an increased proliferation rate owing to loss of cell cycle control and a higher genomic instability secondary to loss of DNA repair.

p53 also appears to be involved in the inhibition of angiogenesis, affecting tumor spread and invasion. Therefore p53 would also be expected to be indicative of the tumor-inherent biological aggressiveness determining the outcome and the prognosis, independent of therapy (or in the absence of systemic therapy). Because p53 may interact positively or negatively with different systemic therapies, the pure prognostic value of p53 can be assessed only by analysis of untreated patient cohorts. The majority of prognostic results derive from retrospective studies, and most of them contained patients who received one or more systemic therapies.

Nonetheless, several studies have tried to address p53 as a prognostic factor (Table 3; [5,11,16,46,48-53])1996 mutations in the p53 gene were reported to constitute the single most important indicator for recurrence and death in breast cancer (45). Blaszyk reviewed 14 studies published between 1993 and 1998, correlating p53 gene mutation and prognosis in breast cancer. Eleven of theses studies used screening methods for p53 analysis. Overall the relative risks reported by the authors were between 2.2 and 3.3 (46).

Elledge reviewed a large number of studies relating the p53 status to prognosis in breast cancer (16). In these 57 studies published from 1992 to 1997, different methods for p53 analysis were used: IHC in 43, screening methods (SSCP, CDGE) in 12, and sequencing in 2 studies only. The authors state that a positive p53 IHC appeared to be consistently associated with other poor prognostic factors such as estrogen and progesterone receptor negativity, a poor grade, and a high proliferative fraction, and was not associated with size or nodal status. The authors concluded that positive p53IHC appeared to be independently associated with a worse prognosis and increases the relative risk of relapse by 1.5. Overall, the authors considered the risk difference to be too small to decide on adjuvant therapy based on the p53 IHC alone.

6. CONCLUSIONS

The p53 gene is one of the most intensively studied molecules in the human genome. From a biologic aspect, p53 is an extraordinary molecule. It has led many researchers to believe that probably every cancer cell has to inherit a defect in the p53 pathway. Thousands of changes may contribute to the malignant phenotype (54). In any case, p53 appears to be involved in a plethora of cell-inherent systems determining cellular proliferation and

Author (ref.)	Journal	No. of patients	No. of patients Technique	Technical specifics	Untreated control	Adjuvant therapy	Prospective for marker question	Prospective for marker Multivariate question analysis	DFS	OS	Authors' conclusions
Elledge (16)	Breast Cancer	694	IHC	PAb 1801+ PAb240	Yes	TAM	No	Independent 0.008	1	0.056	0.056 Prognostic but weak
Beenken (48)	Res Treat 1998 Ann Surg 2001	06	IHC	DACO	Yes	CMF	No	Independent		0.011	Coexpression of <i>p53</i> and <i>c-erh</i> -R7 has
											even more prognostic significance than T or N stage
Balszyk (46)	Int J Cancer 2000	06	Screening	-		CMF or CAF +/-	Yes	Independent 0.0032 0.02	0.0032 0.02	0.0001 0.003	Gene mutations associated with
			IHC	Pab1801+ Pab240+ Pab421		TAM					adverse outcome
Elkhuizen (49) JCO 2000	<i>JCO</i> 2000	361	IHC	Mab DO7	Yes	FU, Doxo, Cyclophosph	No	Independent 0.05	0.05		Associated with higher local relapse rate
Degeorges (50) BCRT 1998	BCRT 1998	282	IHC	Mab DO7	Yes	CMF	No	ns	su	su	
Geisler (11)	Cancer Res 2001	91	Screening	TTGE + sequencing+ Mab DO7	No	doxorubicin	Yes	us	0.0002	0.0034	
Thor (52)	JNCI 1998	595	IHC		No	CAF	No	Independent 0.035	0.035	0.023	
Takahashi (53) IJC 2000	<i>IJC</i> 2000	76	Screening	Screening Yeast functional	No	5 FU + TAM	No	Independent	su	0.032	p53 mutation +
				assay							loss of estrogen receptor affect
											prognosis
Dublin (5)	IJC 1997	277	IHC	Mab D01	Yes	CMF	No	su	ns	ns	

F ٢ 1.1. F ÷ Table 3 . F F ¢ 2 design are only incompletely met. Methods for *p53* analysis do not include sequencing and antibodies used for IHC and screening methods are different as well as the treatment regimens. DFS, Disease-free survival; OS, overall survival. ns, not significant.

death. During cancer development, tumor cells select against the tumorsuppressor functions of p53. Disruption of p53-dependent apoptosis confers a selective advantage during carcinogenesis.

Considering the biologic rationale, p53 appears to be an ideal prognostic and predictive factor. However, more than 10 yr of p53 research have not resulted in a reliable demonstration of the true prognostic and predictive value of p53 and its clinical usefulness in breast cancer or other malignancies. Reviewing the plethora of clinical studies, which are not comparable in many respects, preventing meta-analysis, it would appear that future studies should be planned more carefully to promote the p53 story.

- To assess the true predictive value of *p53* (or any other predictive marker) an *optimal study design* is essential: ideally, *p53* should be analyzed on the basis of monotherapeutic neoadjuvant studies, as the parameter of interest is *response* to treatment with the agent of interest.
- Careful attention is required to *the optimal analysis method*. Any new method to be introduced has to be compared in carefully planned studies to sequencing, which remains the gold standard method.
- As the marker information is needed prior to therapy, it has to be considered that *p53* has to be assessable on biopsy.

Nonetheless, clinical application of p53 status as a criterion for treatment decisions and selection of appropriate therapy could potentially help the clinician to increase response rates to chemo- and radiation therapy, and subsequently increase the rate of radical resection. In addition, patients will be spared the discomfort of ineffective treatment as well as unpleasant side effects.

After all these years of clinical p53 research the final statement has not yet changed: To implement p53 in clinical cancer treatment, well-designed prospective randomized neoadjuvant studies using carefully selected methods of p53 analysis are urgently needed.

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V PREDICTIVE BIOMARKERS IN BODY FLUIDS

12 Occult Metastatic Cells in Breast Cancer Patients

State of the Art, Pitfalls, and Promises

Stephan Braun, MD, Julia Seeber, MD, and Christian Marth, MD

CONTENTS

INTRODUCTION STATE OF THE ART AND PITFALLS IN TUMOR CELL DETECTION CLINICAL RELEVANCE OF TUMOR CELL DETECTION CONCLUSIONS AND PERSPECTIVES REFERENCES

SUMMARY

Blood-borne distant metastasis is the leading cause of cancerrelated death in patients with breast cancer. The onset of this fundamental process can now be assessed in cancer patients using immunocytochemical and molecular assays able to detect even single metastatic cells. However, careful validation of technically confounding variables, including choice of detection antibody, preparation of cellular samples, and size of analyzed sample volumes, is mandatory for reproducible and comparable results. In studies with validated assays, analyses of bone marrow samples show that disseminated cells are present in 20–40% of primary breast cancer patients without any

Cancer Drug Discovery and Development: Biomarkers in Breast Cancer: Molecular Diagnostics for Predicting and Monitoring Therapeutic Effect Edited by: G. Gasparini and D. F. Hayes © Humana Press Inc., Totowa, NJ clinical or histopathological signs of metastasis. The common homing of circulating breast cancer cells in bone marrow is indicative of systemic tumor cell spread and growth of overt metastases in relevant organ sites such as bone, lung, or liver. Recent clinical studies involving more than 3000 breast cancer patients demonstrated that presence of tumor cells in bone marrow at primary diagnosis is an independent prognostic factor for unfavorable clinical outcome. To date, sampling of bone marrow, however, is not a routine procedure in clinical management of breast cancer patients. In this chapter, we review the existing tumor cell assays and discuss their current clinical relevance and perspectives for the clinical management of breast cancer patients.

Key Words: Breast cancer; disseminated tumor cells; circulating tumor cells; prevalence; prognosis.

INTRODUCTION

In breast cancer, recent guidelines for adjuvant systemic therapy result in treatment recommendations for more than 90% of patients even in case of a negative lymph-node status (1,2). The risk of tumor relapse in these patients is considered high enough to recommend adjuvant therapy, even though up to 70% of early stage breast cancer patients are cured by locoregional surgery alone. Therefore, the availability of additional factors enabling individual risk assessment would be desirable to improve identification of patients at risk for relapse.

Although the presence of lymph-node metastasis is a negative prognostic factor for breast cancer and other cancers, it is still not possible to reliably identify those patients who will eventually relapse with metastatic disease only by their lymph node status at primary therapy, indicating that other ways of metastatic tumor cell spread also play an important role. Gene expression profiling (3) and clinical studies (4,5) further indicate that during early stages of breast cancer tumors are more likely to disseminate either via the lymphatic or the hematogenous route rather than simultaneously via both routes.

Advances in the development of immunocytochemical and molecular assays now enable specific detection of metastatic tumor cells even at the single cell stage and thus allow to address the important question of systemic tumor cell dissemination as one of the first crucial steps in the metastatic cascade. Using these technologies, it has become evident that 20–40% of patients with breast cancer harbor occult metastatic cells in their bone marrow even in the absence of any lymph node metastases (stage N0) and clinical signs of overt distant metastases (stage M0). However, these techniques for detection of metastatic cells are not yet established in clinical routine practice, and no international consensus has been reached to recommend a single standardized protocol as benchmark technology.

Several studies on breast cancer patients suggested that presence of disseminated tumor cells (DTCs) in bone marrow represents an additional clinical marker that may be capable to identify those patients who are cured by surgery alone (in the absence of such cells) or may require more specific therapy (in the presence of such cells). One of the intriguing opportunities of this marker might therefore be its use for clinical decision making in risk adapted adjuvant treatment strategies. Another important and (in comparison to other markers) unique application might be the monitoring of therapeutic efficacy in the adjuvant setting with no measurable disease. In this overview, we discuss usefulness and clinical relevance of immunologic and molecular analyses applied for diagnosis of DTCs in bone marrow of breast cancer patients.

2. STATE OF THE ART AND PITFALLS IN TUMOR CELL DETECTION

2.1. Histopathology

Using conventional histopathologic techniques at time of primary diagnosis without clinical signs of metastatic bone disease, the likelihood of identifying DTCs in bone marrow is as low as 4% (6,7). These findings suggest that histopathologic evaluation might not be sensitive enough for the indicated purpose.

2.2. Immunocytochemistry

2.2.1. TUMOR CELLS IN BONE MARROW

To date, most experience with bone marrow screening for DTCs exists for immunocytochemical analyses (Table 1). Numerous studies reported a strong prognostic impact of the presence of DTCs (4,5,8-14), while other investigations failed to do so (Table 2) (15–22). One reason for the discrepant results of clinical follow-up studies is a substantial methodological variation (e.g., sensitivity and specificity of detection antibody, lower detection rate of bone marrow biopsy as compared to bone marrow aspiration, considerable variation in the number of cells analyzed) resulting in a wide range of detection rates between study populations (Tables 1 and 2). Nevertheless, the six most recent studies (5,8-10,12,23) comprising more than 3000 patients consistently reported that the presence of DTCs in bone marrow has a strong prognostic impact on patient survival (Table 2). However, even in these studies, at least three confounding technical factors varied considerably: (1) consistent and blinded analysis of noncarcinoma control patients, (2) diversity of antibodies used for identification of epithelial cells in bone marrow, and (3) number of cells analyzed per patient sample.

Variables		No. of studies (%)
Technical type		
Tissue collection	Bone marrow biopsy	5 (24)
	Bone marrow aspiration	16 (76)
Assay	Immunocytochemistry	17 (81)
2	RT-PCR	4 (19)
Antigens	Mucin-like antigens	5 (24)
e	Cytokeratin	11 (52)
	Mixed	5 (24)
Quantitation of tumor cells	Yes (cytospin)	6 (29)
-	No (biopsy)	5 (24)
	No (cell smears)	6 (29)
	No (RT-PCR)	4 (19)
Detection rates	<10%	2 (9)
	10–19%	4 (19)
	20-29%	4 (19)
	30-40%	6 (29)
	>40%	5 (24)
Clinical type		
Investigated subgroups	All subgroups	20 (95)
	Node-negative, only	1 (5)
	Node-positive, only	
Study type	Mono-center	20 (95)
	Oligo-/multicenter	1 (5)
	Controlled clinical trial	—
	Prospective prevalence study	21 (100)
Size of study population	≤100	8 (38)
	101-500	9 (43)
	>500	4 (19)

Table 1

Technical and Clinical Variability in Bone Marrow Micrometastasis Studies^a

^aData from refs. 4,5,8–15,17–23,78,80–82.

Pantel et al. (24) designed a study to evaluate some of the variables affecting the immunocytochemical detection of individual epithelial tumor cells in bone marrow. Bone marrow aspirates were taken from 368 patients with primary carcinomas of the breast, lung, prostate, or colon–rectum using an alkaline phosphatase antialkaline phosphatase staining technique. DTCs were detected with specificity proven monoclonal antibodies CK2 and

A45-B/B3. In the first line, detection rate was affected by blood contamination of the aspirates, the number of aspirates analyzed, and the number of marrow cells screened per aspiration site. The significantly lower detection rate by monospecific anti-CK-18 antibody CK2 as compared to antibody A45-B/B3, which detects heterodimers CK-8/18 and CK8/19, indicated that assay sensitivity may, however, be further affected by factors of the microenvironment or altered gene expression during metastasis. Several recent studies analyzing gene expression (*3*) and protein expression (*25,26*) indicated that down-regulation of single CK peptides (e.g., CK-18) may be necessary to alter cell plasticity in order to increase invasiveness and metastasis. Thus, beyond mere basic technical aspects (e.g., sensitivity and specificity), tumor-related processes significantly influence assay validity.

Sufficient methodological validation of the detection antibodies has only been reported for anticytokeratin antibodies (7,24,27-29). The rare occurrence of single CK-positive cells in aspirates of noncarcinoma control patients (9) points to minimal-yet acceptable-technical variations within a biological system. The resulting risk of false-positive findings in cancer patients can be minimized by using morphological criteria in addition to immunostaining for diagnosis of DTCs (27,28). Additional justification for using cytokeratin-specific antibodies in screening assays for occult breast carcinoma cells can be derived from several recent studies, describing phenotypic characteristics of CK-positive DTCs as similar to those usually found in malignant solid tumors (30-33). Furthermore, in single CK-positive DTCs, numerical chromosome aberrations were detected by interphase fluorescent *in situ* hybridization analysis (34); multiple mutations by whole-genome amplification (35); and multiple chromosomal aberrations by combined transcriptome and genome analysis of single micrometastatic cells (36-38).

Both the International Society of Cell Therapy (ISCT) and the National Cancer Institute (NCI) have recognized the need for standardization of the immunocytochemical assay and for its evaluation in prospective studies (27,39). On the basis of data available from published methodological analyses (9,24,27,29) such a standardized assay may consist of a specificity-proven, anti-CK monoclonal antibody (i.e., A45–B/B3) and a sufficient sample size (i.e., 2×10^6 mononucleated cells per patient) obtained from two aspiration sites. The use of new automated devices for the microscopic screening of immunostained slides may help to read slides more rapidly and to increase reproducibility of the read-out process (40-42).

2.2.2. TUMOR CELLS IN PERIPHERAL BLOOD

Peripheral blood would be an ideal source for the detection of disseminated tumor cells because of an easy sampling procedure. The presence of malignant cells in peripheral blood was described several decades ago

Authors (ref.)	Antigens	Preparation	Technique	Technique No. of patients	Detection rate (%)	Prognostic value
Bone marrow						
Porro et al. (22)	Mucin	Biopsy	ICC	159	16	None
Salvadori et al. (21)	Mucin	Biopsy	ICC	121	17	None
Mathieu et al. (20)	Mucin/CK	Biopsy	ICC	93	1	None
Courtemanche et al. (18)	Mucin	Biopsy	ICC	50	8	None
Singletary et al. (19)	Mucin/CK	Cell smears	ICC	71	38	None
Cote et al. (14)	Mucin/CK	Cell smears	ICC	49	37	DFS, OS
<u> </u>	Mucin/CK	Cell smears	ICC	100	38	DFS^a , OS^a
Diel et al. (12)	Mucin	Cell smears	ICC	727	43	DFS^a , OS^a
	CK	Cytospins	ICC	234	38	n.d.
	CK	Biopsy	ICC	128	19	DFS^a , OS^a
	Mucin	Cell smears	ICC	350	25	DFS, OS
	CK	Cytospins	ICC	581	28	None
Braun et al. (9)	CK	Cytospins	ICC	552	36	$DDFS^{a}, OS^{a}$
Braun et al. (4)	CK	Cytospins	ICC	150	29	DDFS, OS^a
Gerber et al. (5)	CK	Cytospins	ICC	484	31	DFS^{a} , OS^{a}
Gebauer et al. (8)	Mucin/CK	Cell smears	ICC	393	42	DFS^a , OS
Wiedswang et al. (23)	CK	Cytospins	ICC	817	13	DDFS ^a , BCSS ^a
Datta et al. (82)	CK19	Cell suspension	RT-PCR	34	26	DFS
Fields et al. (81)	CK10	Cell suspension	RT-PCR	83	71	DEC

Table 2

Vannucchi et al. (80)	CK19	Cell suspension	RT-PCR	33	48	DFS
Slade et al. (78)	CK19	Cell suspension	RT-PCR	23	61	None
Peripheral blood						
86)	CK19, EGF-R	CK19, EGF-R Cell suspension	RT-PCR	21	81	None
		Cytospins		$22 (133)^b$	42	na
Smith et al. (59)		Cell suspension		$22 (145)^b$	50	na
Stathopoulou et al. (57)	CK 19	Cell suspension	RT-PCR	148	30	DFS^a , OS^a
		Cell suspension		114	25	None
Racila et al. $(7I)$		Cell suspension		30	76	na
Witzig et al. (106)	CK	Cytospins	ICC	75	28	na
						9 10 11

BCSS, Breast cancer specific survival; CK, cytokeratin; DFS, disease-free survival; DDFS, distant disease-free survival; FC, flow cytometry; H&E, hematoxylin and eosin staining; hMAM, human mammaglobin; ICC, immunocytochemistry; IHC, immunohistochemistry; OS, overall survival; RT-PCR, reverse-transcriptase polymerase chain reaction; na, not analyzed.

"Prognostic value supported by multivariate analysis.

^bNumber of samples analyzed.

(43,44), and concerns toward intraoperative manipulation of the tumor causing uncontrolled tumor shedding were supported by studies on gastrointestinal and prostate cancer patients (45-50). More recent reports have confirmed the malignant nature of the detected cells using cytogenetic or molecular analysis (51,52). From model systems, it was estimated that about 10⁶ tumor cells/g of tumor tissue are shed daily into the blood, although such model calculations might overestimate the number actually shed in vivo (53-55). Yet, for circulating tumor cells (CTCs), blood is only a temporary compartment in which no decision on the subsequent fate of CTCs per se is made, except for trapping of tumor emboli in the first capillary bed these cells encounter. Yet despite the fact that most cells that once entered the blood circulation survive the blood passage, metastasis remains an inefficient process with successful onset of micrometastasis by only a limited number of cells (56). In consequence, from currently available studies it remains unknown, which fraction of CTCs survive and inherit the potential for metastatic growth in the new secondary microenvironment. This ignorance, which in part is due to the nature of the analysis of CTCs, hence probably less clonally selected, and in part to an enormous discrepancy between the actual circulating tumor load and the fraction available for analysis, explains the undetermined clinical value of presence of CTCs (Table 2). In breast cancer, few relatively small studies with different technical approaches suggested that presence of CTCs correlated with stage and course of the disease (57-59). For other solid tumors, several reports have also shown a prognostic impact of CTCs in blood (60-64). However, it remains debatable if these studies are empowered enough to answer the question of prognostic impact with a reasonably low β -type error, in view of relatively small study populations and sole focus on *p*-values for α -type error.

2.3. Specific Tumor Cell Enrichment and Depletion

Some of the discordant results from studies examining the role of DTCs and CTCs might be also due to the low frequency of these cells, and hence the technique-related variable degree of the sampling error. New enrichment techniques based on improved density gradient methods (65) and immunomagnetic procedures (66,67) may increase assay sensitivity in parallel reducing the degree of sampling error, which in consequence might help to increase data consistency of the clinical significance of DTCs and CTCs. Immunomagnetic enrichment utilizes differential expression of either tumor- or tissue-specific antigens by tumor cells as compared to mesenchymal cells in bone marrow and peripheral blood. For positive separation of tumor cells, beads coated with tumor- or tissue-specific antigens select DTCs and CTCs, while for negative separation, antibodies directed against hematopoietic or mesenchymal markers are coated to ferromagnetic beads (67-72). These selection strategies have the additional advantage that

the tumor cells are still viable and can be used for additional studies including the propagation of malignant cells in vitro (73). At present, these new techniques are costly, time consuming, and their benefit over "standard" Ficoll gradients remains to be substantiated in further studies.

2.4. Genetic and Epigenetic-Based Assays

The lack of a unique genetic marker for solid cancers such as breast cancer requires sufficient sensitivity and specificity of tumor-cell associated genetic markers used for detection of DTCs and CTCs within a body compartment. The major limiting factor in detection of DTCs and CTCs by reverse transcriptase polymerase chain reaction (RT-PCR) is illegitimate transcription of tumor-associated or epithelial-specific genes reported for mesenchymal cells (74,75). Moreover, owing to the extreme genetic instability of breast carcinoma cells, deficient expression of the marker gene in DTCs and CTCs may lower the actual sensitivity in vivo as compared to experimental model systems that use tumor cell lines to estimate assay sensitivity. Quantitative real-time PCR provides interesting prospects for better quantification of the tumor cell load, provided specific markers are available. The studies conducted so far in bone marrow and peripheral blood (Table 1) applied CK-19, mucin, and human mammaglobin (hMAM) mRNA markers (57,59,76-82) yet with uncertain specificity (57,75,83-87). To date, the marker most extensively investigated for the detection of tumor cell (i.e., DTC/CTC) derived transcripts is CK-19. Using nested RT-PCR or competitive quantitative RT-PCR, with the number of CK-19 transcripts being normalized to the number of (breast cancer unrelated) ABL transcripts, several investigators found CK-19 transcripts in bone marrow and peripheral blood of early or metastatic breast cancer patients in 63–74% and 30–52%, respectively (57,59,78). In establishing a quantitative realtime RT-PCR assay, however, we encountered serious difficulties being unable to obtain discernible crossing-points for samples from 129 cancer and 16 control patients (83), strongly suggesting that CK-19 transcripts in BM—to some extent—are not of epithelial/tumor origin. Our assay used CK-19 primers with known absence of reactivity with CK-19a/CK-19b pseudogenes in lymphoid tissues (88). These real-time RT-PCR data essentially confirm our previous results using nested RT-PCR. While in all breast cancer tissues hMAM, EGF-R, and CK-19 transcripts were detectable, blood samples from 31 healthy controls and 20 patients with hematological malignancies revealed absence of hMAM transcripts but presence of CK-19 transcripts in 27% and EGF-R transcripts in 10% of these cases (84). These data are in contrast to the above cited studies on prevalence and assumed clinical relevance of CK-19 transcript detection in peripheral blood and bone marrow. The data on the limited value of currently propagated RT-PCR assays for the detection of DTC and CTC-derived transcripts point to the urgent need of improved approaches and studies demonstrating the actual value.

Beyond genetic characteristics, with the prevailing uncertainty about their specificity in solid tumors, changes in the status of DNA methylation, known as epigenetic alterations, are among the most common molecular alterations in human neoplasia (89), including breast cancer (90). Cytosine methylation occurs after DNA synthesis by enzymatic transfer of a methyl group from the methyl donor S-adenosylmethionine to the carbon-5 position of cytosine. Cytosines (C) are methylated in the human genome mostly when located 5' to a guanosine (G). Regions with a high G:C content are so-called CpG islands. It has been increasingly recognized over the past 4-5 yr that the CpG islands of many genes, which are mostly unmethylated in normal tissue, are methylated to various degrees in human cancers, thus representing tumor-specific alterations (89,91). In a recent study on the prognostic value of DNA methylation in serum of breast cancer patients, methylated serum DNA for RASSF1A and/or APC was found to discriminate significantly and independently of established clinical factors between patients with more favorable (unmethylated serum DNA) and those with poor prognosis (methylated serum DNA) (92). The tumor specificity and the possibility to use blood tests, together with the option for specific therapy using demethylating agents (93), suggest the potential of this approach for diagnosis and monitoring patients' course of the disease. Yet, it still remains to be shown whether the assessment of epigenetic alterations and their specific treatment play the assumed functional role that determines metastatic growth, and hence the fate of the patient.

3. CLINICAL RELEVANCE OF TUMOR CELL DETECTION

So far, studies on the prognostic value of DTCs in bone marrow are almost exclusively based on immunocytochemical data (Table 1). An important initial question was whether the presence of epithelial antigen-positive cells is correlated with established risk factors in breast cancer, such as tumor size or lymph node involvement. Diel et al. (12) found a significant correlation between bone marrow positivity, as assessed by antitumor-associated glycoprotein-12 (TAG-12) immunocytochemistry, and tumor size, nodal status, histopathologic tumor grading, as well as postmenopausal status. The London Ludwig Cancer Institute Group described that presence of EMA-positive cells in bone marrow was significantly related to lymph node involvement, peritumoral vascular invasion, and primary tumor size (94). Studies using different anticytokeratin monoclonal antibodies demonstrated merely a tendency toward correlation between detection of cytokeratin-positive cells in bone marrow and locoregional lymph node involvement (7,95). Applying the broad-spectrum anticytokeratin monoclonal antibody A45-B/B3 for DTC detection (9), a significant association of such cells in bone marrow with the diagnosis of inflammatory breast cancer, tumor size, extensive lymph node metastasis of >10 nodes, and tumor grade was reported.

To assess the clinical significance of DTCs in bone marrow, follow-up studies were initiated. Using a polyclonal EMA antibody, Mansi et al. (10) detected metastatic cells in 25% of bone marrow samples. In their 6-yr follow-up analysis, univariate statistics revealed that the immunocytochemical finding predicts for an increased rate of relapse in bone and other distant sites, as well as decreased overall survival. Multivariate analysis both after 6 (96) and more than 12 yr of clinical follow-up (10) indicated that the prognostic impact of EMA-positive cells was not independent of established risk factors, such as tumor size, grade, and lymph node status. Using another mucin marker TAG-12, Diel et al. (12) reported a detection rate of 43% in a collective of 727 primary breast cancer patients. After a median follow-up time of 36 mo (3-108 mo), the presence of TAG-12-positive cells was described as being superior to axillary lymph node status, tumor stage, and tumor grade as an independent prognostic indicator for both metastasisfree and overall survival. But in contrast Gebauer et al. (8) detected metastatic cells in 42% of 393 breast cancer patients. Follow-up data were available for up to 10 yr after surgery. The multivariate analysis revealed independent prognostic relevance for bone marrow status, although tumor size and axillary lymph node status were superior prognostic factors.

Using different cocktails of monoclonal antibodies to cell-surface antigens and cytokeratins, Cote et al. (14), Harbeck et al. (13), and Gebauer et al. (8) were able to derive prognostic information from the presence of occult metastatic cells in bone marrow. A potential disadvantage of some of the earlier studies is substantial methodological variation. Therefore, a prospective two-center study in 552 primary breast cancer patients was initiated, using a validated immunoassay (24,29). The test results could be reproduced independently at both study centers (9). In this study, multivariate regression analysis verified that the presence of DTCs in bone marrow predicts poor prognosis overall (Fig. 1) and independent of the lymph node status (Fig. 2). These findings demonstrate for the first time that by using an identical assay for detection of DTC, reproducible results can be generated. In parallel to our study, a Norwegian study was launched and recently published, as the currently largest available single center study (817 patients), with a median follow-up of 49 mo (23). The independent prognostic value of DTC in bone marrow was clearly confirmed for distant metastasis-free and breast cancer-specific survival. In combination of independent prognostic factors, such as tumor size, grading, and lymph node status, they were able to classify subgroups of both node-negative and node-positive patients

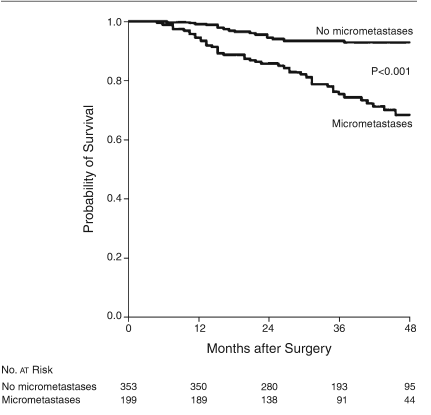


Fig. 1. Overall survival of 552 breast cancer patients according to presence or absence of DTC in bone marrow, after a median observation time of 38 mo (log-rank test). (Adapted from ref. 9.)

into excellent and high-risk prognosis groups (Fig. 2), supporting the usefulness of DTC screening for patient stratification.

4. CONCLUSIONS AND PERSPECTIVES

Various immunocytochemical and molecular methods have been applied to detect occult hematogenous tumor cell spread in breast cancer patients. Numerous studies have marked the prognostic implications associated with the presence of occult metastatic cells (OMCs) at the time of diagnosis. Yet, screening of bone marrow micrometastasis is currently not accepted as a routine diagnostic tool. Reasons for this are the lack of international consensus and consequently the lack of studies demonstrating both improved risk stratification and patient outcome in the setting of clinical trials. International consensus is now urgently needed regarding quality control issues

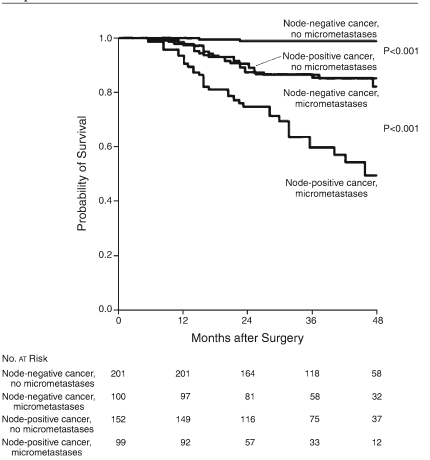


Fig. 2. Overall survival of 552 breast cancer patients according to presence or absence of DTC in bone marrow and lymph-node status, after a median observation time of 38 mo (log-rank test). (Adapted from ref. 9.)

and criteria for acceptable technical assay performance, such as false-negative and false-positive rates, for clinically applicable assays to permit comparisons between different assay platforms. Finally, to support efforts on clinical trial design, marker implementation into current risk classification systems, such as the Tumor–Node–Metastasis (TNM) classification system is needed. A useful proposal has recently been made by the International Union Against Cancer (UICC) (97). The most recent TNM classification for breast cancer (98) does not qualify the presence of single cancer cells in peripheral blood or bone marrow as metastasis (stage M0), but it optionally reports the presence of such cells together with their detection method (e.g., M0[i+] or [mol+]).

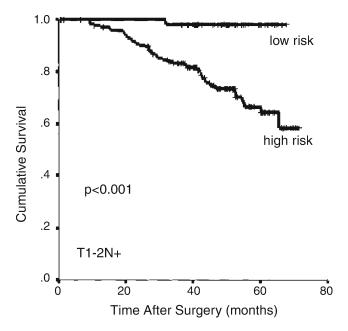


Fig. 3. Overall survival of breast cancer patients according to presence or absence of independent prognostic factors, after a median observation time of 49 mo (log-rank test). Low-risk profile (tumor size <2 cm, estrogen/progesterone receptor-positive and bone marrow-negative) vs high-risk profile (tumor size >2 cm and/or estrogen/progesterone receptor-negative and/or bone marrow-positive). (Adapted from ref. 23.)

Beyond merely adding another prognostic factor to the plethora of such markers in breast cancer, it needs to be emphasized that assessment of occult hematogenous tumor cell spread inherits the potential for a tool for prediction and monitoring of efficacy of systemic therapy (99–103). In contrast to lymph nodes, which are generally accepted as "indirect" marker of hematogenous tumor cell spread and, hence, risk of systemic spread but which are also generally removed at primary surgery and unavailable for follow-up evaluations, bone marrow and blood can be obtained repeatedly in the postoperative course of the patient. In Fig. 3, for example, the only prognostic factor available for follow-up risk assessment is the presence of bone marrow micrometastasis. The clinical value of such examinations has been strongly suggested by clinical studies on a total of almost 500 patients, which identified the prognostic relevance of OMC present in bone marrow several months after diagnosis or treatment when no relapse has occurred until that date (100, 104, 105). The potential of a surrogate marker assay that permits immediate assessment of therapy-induced cytotoxic effects on occult metastatic cells is therefore evident, as indicated previously (100). Since repeated bone marrow sampling might not be easily implemented into clinical study protocols for breast cancer, serial examinations of blood for CTCs or tumor cell-associated nucleic acids might be more acceptable for most patients and clinical investigators than repeated bone marrow aspirations. The detection and characterization of CTCs in peripheral blood of cancer patients has therefore received much attention in recent years and could lead to strategies for evaluation of therapeutic efficacy. The availability of a standardized, reliable blood test could enable implementation of CTCs as a surrogate marker for clinical development of new anticancer agents and optimization of existing treatment protocols. For the time being and with validated assays being available for the evaluation of bone marrow only, prospective clinical studies are now required to evaluate whether eradication of DTCs in bone marrow after systemic therapy translates into a longer disease-free period and overall survival.

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13

Circulating HER2/neu

Clinical Utility

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CONTENTS

INTRODUCTION METHODS OF DETERMINING HER2/NEU STATUS PREVALENCE OF CIRCULATING HER2/NEU LEVELS IN BREAST CANCER SERUM HER2/NEU AS A PROGNOSTIC INDICATOR SERUM HER2/NEU AS A PREDICTIVE INDICATOR MONITORING HER2/NEU ECD LEVELS IN MBC PATIENTS TREATED WITH HORMONE OR CHEMOTHERAPY MONITORING METASTATIC BREAST CANCER PATIENTS TREATED WITH HERCEPTIN-BASED THERAPIES SERUM HER2/NEU LEVELS AND DETECTION OF EARLY RECURRENCE **CONCLUSIONS** REFERENCES

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SUMMARY

The HER2/neu oncogene and its p185 receptor protein is an indicator of a more aggressive form of breast cancer. The HER2/neu status guides Herceptin[®] therapy, specifically directed to the extracellular domain (ECD) of the HER2/neu oncoprotein. The HER2/neu ECD is shed from cancer cells into the circulation and is measurable by immunoassay. We performed an in-depth review of the peer-reviewed literature on circulating ECD levels with respect to prevalence, prognosis, prediction of response to therapy, and monitoring of breast cancer. Studies showed that the prevalence of an elevated ECD in patients with primary breast cancer varied between 0% and 38% (mean 18.5%) while in metastatic breast cancer the range was from 23% to 80% (mean 43%). Some women who have HER2/neu-negative tumors by tissue testing develop elevated ECD levels in metastatic disease. Elevated ECD levels have been correlated with indicators of poor prognosis, for example, overall survival and disease-free survival. Elevated ECD levels predict a poor response to hormone therapy and some chemotherapy regimens but can predict improved response to combinations of Herceptin and chemotherapy. Many studies support the value of monitoring ECD levels during breast cancer progression, as serial increases precede the appearance of metastases using imaging techniques, and longitudinal ECD changes paralleled the clinical course of disease. The monitoring of circulating HER2/neu ECD levels provides a tool for assessing prognosis, predicting the response to therapy and for earlier detection of disease progression and intervention with appropriate therapy. Additional prospective studies are required to validate these potential applications.

Key Words: HER2/neu; extracellular domain; oncoprotein; breast cancer.

1. INTRODUCTION

The activation and overexpression of cellular oncogenes has been considered to play an important role in the development of human cancer (1). An important member of the oncogene family is the growth factor receptor known as the <u>H</u>uman <u>Epidermal</u> Growth Factor <u>Receptor-2</u> (HER2) (2), which is also referred to as *HER2/neu* or c-*erb*-B2. *HER2/neu* is structurally and functionally related to the v-*erb*-B retroviral oncogene (3) and is part of the *HER* family which also includes HER1 or EGFR (epidermal growth factor receptor) and *HER3* and *HER4* (4).

The *HER2/neu* oncogene has been localized to chromosome 17q and encodes a transmembrane tyrosine kinase growth factor receptor that is expressed on cells of epithelial origin. The full-length glycoprotein has a

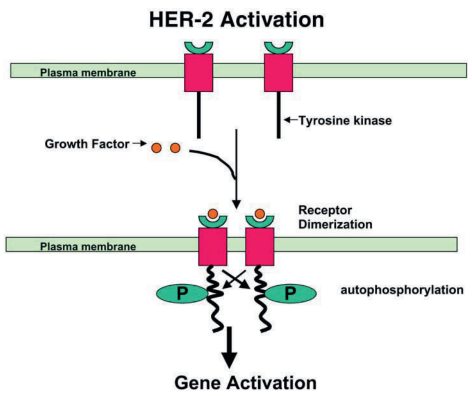


Fig. 1. HER2/neu signaling pathways are activated by homo- and heterodimerizations with extracellular domains of the other HER family members.

molecular weight of 185,000 daltons (p185) and is composed of the internal tyrosine kinase domain, a short transmembrane portion, and has an extracellular domain (ECD) similar to the three other members of the *HER* family (2,5). The ECD portion of the receptor protein is heavily glycosylated, has a molecular weight in the 97–115-Kda range and has been shown to be shed into cultured fluids of SKBR-3 cells (6), as well as plasma (7) and serum (8,9) from normal individuals and patients with breast cancer (7–9).

The mechanism of activation of the HER2/neu pathway is not completely understood but studies have shown that the ECDs of the *HER* family of receptor tyrosine kinases form homodimers and heterodimers and that receptor dimerization activates a cascade of events in the HER-2/neu signaling pathway (4). A ligand that binds to the HER2/neu receptor has not been identified but a family of peptide ligands named *neu* differentiation factors or heregulins have been identified that bind to the HER3 and HER4 receptors, inducing heterodimerization with the HER2/neu receptor and therefore inducing transphosphorylation and thus activation of the HER2/neu receptor (Fig. 1). Heterodimers between the EGFR and HER2/neu can also form upon binding of a ligand such as epidermal growth factor or transforming growth factor- α to the EGF receptor (10). Dimerization of the receptor results in tyrosine phosphorylation of the HER2/neu kinase with subsequent activation of downstream transduction pathways and signaling through ras, c-Src, phosphatidylinosital 3 kinase (PI3K), and phospholipase C (PLC)- γ pathways. Formation of heterodimers increases the affinity of the partnering receptor for its ligand and results in potentiation of the mitogenic signal (11).

In a report by Codony-Servat et al. (12), it was demonstrated that cleavage of the HER2/neu ECD involves metalloproteinase (MMP) activity and that the process of cleavage was inhibited by the MMP inhibitor TIMP-1 but not by TIMP-2. They also showed that HER-2/neu ECD shedding was inhibited by broad spectrum MMP inhibitors such as EDTA (ethylenediamine tetracetic acid), TAPI-2 and Batimastat. The report also confirmed data from Christianson et al., showing that HER2/neu ECD cleavage results in the release of a truncated membrane-bound phosphorylated p95 fragment (13).

Molina et al., showed in a 2001 report (14) that HER2/neu shedding was activated by 4-aminophenylmercuric acetate (APMA, a well known MMP activator) in HER2/neu overexpressing breast cancer cells and it could be blocked by the MMP inhibitor Batimastat. The increase in ECD shedding also enhanced the production of the p95 fragment. In the same report, Molina et al. (14), demonstrated that Herceptin[®], an anti-HER2/neu therapy, had a direct inhibitory effect on the basal and activated processes involved in HER2/neu cleavage from HER2/neu-overexpressing breast cancer cells. The HER2/neu ECD shedding that was activated by APMA could be blocked with Herceptin, leading to a reduction in the release of the p95 fragment. The analysis of 24 human breast tumors by this group showed that the p95 fragment could be detected in only 14 out of 24 of the specimens, and the p95 band was highly variable. Studies continue to determine the potential clinical value of the phosphorylated membrane fragment (14).

Trastuzumab (commonly referred to as Herceptin, manufactured by Genentech, San Francisco, California, USA) is a humanized monoclonal antibody (MAb) developed to target the HER2/neu receptor that is overexpressed on some cancer cells, including 25–30% of breast cancers as well as a variety of other epithelial cancers. Herceptin binds with high affinity to the ECD of HER2/neu and inhibits proliferation of tumor cells that overexpress HER2/neu protein. The results of a large multicenter phase III clinical trial demonstrated that Herceptin, when added to conventional chemotherapy, can provide benefit to patients with metastatic breast cancer that overexpresses HER2/neu. As compared with the best available standard chemotherapy, concurrent treatment with Herceptin and first line chemotherapy was associated with significantly longer time to disease progression, a higher rate of response, longer duration of response, and improved overall survival (15).

In this chapter, we will review HER2/neu circulating ECD levels in relation to prevalence, prognosis, prediction of response to therapy, monitoring in metastatic breast cancer, and monitoring for early detection of recurrence.

2. METHODS OF DETERMINING HER2/neu STATUS

The most widely accepted method for measuring HER2/neu protein (p185) overexpression is immunohistochemistry (IHC) (16,17) whereas the number of *HER2/neu* gene copies or gene amplification is determined by using a fluorescence *in situ* hybridization (FISH) assay (16,17). Enzyme-linked immunoabsorbent assays (ELISAs) have been used since 1991 to quantitate either the full length p185 in tumor tissue (7) or the soluble circulating HER2/neu ECD in serum (8) or plasma (7).

However, there are limitations to tissue testing, the most important being that they are one-time tests used to determine the HER2/neu status of the primary breast cancer which in turn determines eligibility for Herceptin. Herceptin is given primarily in the metastatic setting, yet the HER2/neu status is determined from the original breast tumor, which may have been removed many years earlier. It is clear from this review of the literature (and will be discussed later) that the HER2/neu status of a breast cancer patient can change from the primary to the metastatic phase.

Tissue testing for HER2/neu protein overexpression by IHC and FISH testing for DNA amplification are subject to technical problems. These include, but are not limited to, differences in the methodology between laboratories and the operator, variability in operator interpretation, and variation in reagents. Although the FISH technology is reproducible, the major drawback is that the FISH equipment is expensive and not widely available in diagnostic pathology laboratories.

The third method used to quantitate full length p185 in tumor tissue or the circulating ECD in serum or plasma is the immunoassay. Zabrecky et al. (6), using MAb directed to the ECD (20), demonstrated that the ECD was shed into the culture supernatant of SK-BR-3 breast cancer cells. Studies employing specific MAbs against the HER2/neu protein and using immuno-precipitation and Western blot techniques showed that the ECD was a gly-coprotein with molecular weight between 97 and 115kDa (6). Subsequent studies illustrated that the ECD could be detected in the plasma of healthy subjects and is elevated in women with primary and metastatic breast cancer (7). These observations were later confirmed by Leitzel et al. (8) and Pupa et al. (9).

In the last several years, many reports have described a variety of ELISA formats that have been used to quantitate the ECD in serum or plasma of breast cancer patients and control groups. However, it has been difficult to compare results between publications owing to a lack of standardization

between the ELISAs. For example, three publications reporting ECD results with one particular commercial assay (Triton-Ciba Corning-Chiron) used at least three different cutoff levels (3 U/mL, 12 U/mL and 30 U/mL) to separate normal and diseased populations (8,21,22). In some reports, antibody specificity or assay validation for HER2/neu has not been demonstrated (23), nor have adequate references been provided to demonstrate that the antibodies in the ELISAs specifically detect the HER2/neu ECD.

In summary, IHC and FISH testing can be used to determine the HER2/neu status in primary tumor tissue but are not adequate for assessing HER2/neu status of a woman after the tumor is removed by surgery. In contrast, the immunoassay method for measuring the circulating HER2/neu ECD is the only way to obtain a real-time status of HER2/neu and the only practical way to monitor changes in the HER2/neu ECD levels.

3. PREVALENCE OF CIRCULATING HER2/neu LEVELS IN BREAST CANCER

We reviewed a total of 55 publications from which data on the prevalence (7,8,22–73) of elevated levels of circulating HER2/neu could be extracted. The data are summarized in Fig. 2 and represent circulating HER2/neu ECD measurements in more than 6500 patients with breast cancer. A review of 24 references used to evaluate ECD levels in primary breast cancer showed that approx 18.5% of the 1923 patients had circulating HER2/neu ECD levels that were above the control cutoff described in each publication.

In contrast, a review of 45 references and 4622 patients with metastatic breast cancer (MBC) showed that 43% of the patients had circulating HER2/neu ECD values above the normal cutoff for the control group. In 15 of the 45 publications ECD levels were elevated above the control group in over 50% of the patients studied.

It is interesting to note that the data in Fig. 2 came from essentially six different assays. The automated Immuno-1 HER2/neu test (manufactured by Bayer Health Care, Tarrytown, NY) and the Oncogene Science manual microtiter plate HER2/neu test (manufactured by Oncogene Science, Cambridge, MA) are essentially the same assays since both use anti-HER2/neu MAbs NB-3 and TA-1 (20), a soluble p97–115-kDa standard, and a 15 ng/mL cutoff (35,65,74). It should also be noted that there was a strong correlation between the automated and manual HER2/neu assays (35,65,74). Both the automated and manual HER2/neu assays have been cleared by the US Food and Drug Administration (FDA) for use in the management and monitoring of women with metastatic breast cancer. These are the only two assays currently cleared for the measurement of the circulating p97–115-kDa HER2/neu ECD. There were 20 publications in Fig. 2 that used the two methods described above. There were 21 references to the Triton/Ciba/Chiron

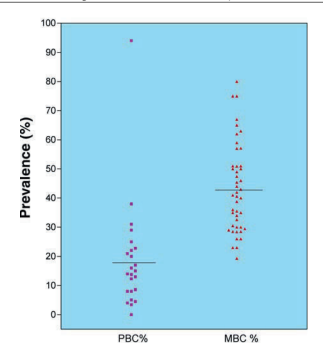


Fig. 2. Circulating HER2/neu ECD measurements in more than 6500 patients with breast cancer. PBC % = % of primary breast cancer patients who had HER2/neu ECD levels above the control cutoff described in each publication (each square = 1 publication; 24 references, n = 1923 patients total, mean = 18.5% above cutoff). MBC % = % of metastatic breast cancer patients who had HER2/neu ECD levels above the control cutoff described in each publication (each triangle = 1 publication; 45 references, n = 4622 patients total, mean = 43% above cutoff).

assays with 11 different cutoffs ranging from 5 to 30 U/mL as well as a 120 fmol/mL or 450 fmol/mL cutoff. These assays are essentially the same and are no longer commercially available. There were five references to the Nicherei assay, three references to the Calbiochem or ORP assay, three references to the Dianova assay, and two references to the Bender assay. We could not find any references that described the antibody specificities or standard material used in the Calbiochem, Dianova or the Bender assays nor could we find any references that validated biochemically that the assays clearly measured the circulating HER2/neu ECD. The Calbiochem, Dianova, and Bender assays are available as research use only assays, which means that their performance characteristics have not been determined. In fact, the Bender assay claims to measure the circulating full-length

p185 nor does the manufacturer present data to support their claim. Several other reports, however, have reproducibly demonstrated that the only HER2/neu fragment found circulating is the truncated p97–115-kDa (6–9) ECD, so it is unclear what the Bender assay actually measures.

In a report by Andersen et al. (25), it was shown that elevated HER2/neu ECD levels were detected in the serum of 8% of preoperative breast cancer patients and in only 3% of postoperative sera from patients without recurrent breast cancer. In contrast, 59% (55 out of 93) of patients with recurrent breast cancer developed elevated HER2/neu ECD levels. It was also reported that elevated ECD levels were detected significantly more often in patients with distant metastases than in patients with recurrent disease restricted to local metastasis (68% vs 19%). This observation was supported by Watanabe et al. (21), who concluded that the circulating HER2/neu ECD level was closely related to tumor mass, since the HER2/neu ECD level in recurrent disease.

In the report by Andersen (25) it was shown that 14 of 24 patients who had immunohistochemistry (IHC)-positive breast tumors also had elevated HER2/neu serum levels during the metastatic phase of the disease. In contrast, 28 of 82 (34%) patients, who had IHC-negative primary breast tumors, developed elevated serum levels during the metastatic phase. Kandl et al. (48) also reported that some patients with negative HER2/neu tumor staining developed extremely high serum levels of HER2/neu during the evolution of metastatic disease which also correlated with extensive MBC. Molina also reported that 23% of patients with recurrent breast cancer with no tissue overexpression had elevated ECD levels, once again supporting the concept that there is a subpopulation of women with HER2/neu–positive tumors that are not identified by tissue testing of the primary breast cancer.

4. SERUM HER2/neu AS A PROGNOSTIC INDICATOR

We analyzed 20 publications and outcomes data from 4430 breast cancer patients (3338 metastatic patients and 1092 primary breast cancer patients) to see if elevated serum HER2/neu ECD levels correlated with poor prognosis. This analysis took into account several indicators of prognosis including time to progression (TTP), overall survival (OS), and disease-free survival (DFS). The data is summarized in Table 1 and in some cases, where available, the hazard ratios had also been calculated and these have been summarized in Fig. 3.

Bewick et al. examined the clinical significance of shed ECD plasma levels in metastatic breast cancer (MBC) patients (27) and showed that 46% of the patients receiving high-dose chemotherapy (HDCT) had elevated ECD levels compared to controls. The results showed that patients with high levels of ECD had a significantly poorer overall survival and progression-

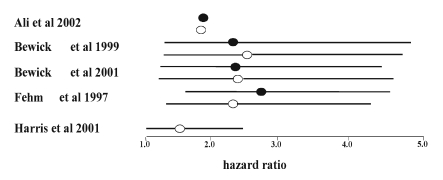


Fig. 3. Summary of the hazard ratios and 95% confidence intervals calculated in the respective publications quoted for serum HER2/neu as a prognostic marker in patients with breast cancer. Overall survival (closed circle) and disease-free survival (open circle).

free interval following high-dose therapy with paclitaxel and autologous stem cell transplantation compared to individuals without elevated ECD levels. For instance, the median overall survival of patients with a low ECD level was 29.8 mo, significantly longer than the 15.9 mo seen in patients with high ECD levels. In addition, the progression-free survival period was significantly longer at 13 mo for patients with low ECD levels than the 8.6 mo observed for patients with high ECD levels (27). In a similar study by Harris et al., patients treated with HDCT and bone marrow transplant who had high ECD levels did worse than patients receiving the same treatment but with low ECD levels (43).

When the prognostic value of this oncoprotein was evaluated by Molina et al., the patients with abnormally high presurgical serum HER2/neu levels had a worse prognosis than those patients with normal levels, in both node-negative and node-positive patients (57). In addition, serum ECD levels in patients with advanced breast cancer were related to the site of recurrence with significantly higher values in patients with metastases (45.4%) than in those with locoregional recurrence (9.2%). In 2002, a report by Ali et al. demonstrated that the median overall survival for MBC patients with normal levels of serum HER2/neu Was 17.1 mo. In contrast, women with normal levels of serum HER2/neu ECD had a median overall survival of 29.0 mo, indicating that elevated HER2/neu levels correlated with poor clinical outcome (24).

In summary, the collective evidence presented in Table 1 from more than 4000 breast cancer patients shows a strong correlation between elevated HER2/neu ECD levels and worse prognosis as demonstrated by a decrease in time to progression, decreased overall survival and decreased disease-free progression.

Table 1 Summary of Data Showing HER2/neu Relationship to Indicators of Prognosis in Patients With Breast Cancer

Authors (ref.)	Patients	Therapy	Prognostic indices (elevated vs nonelevated)	Significance
Ali et al. 2002 (24)	566 MBC	Megestrol or fadrozole; 2nd line	TTP median 89 vs 176 d OS 515 vs 869 d	p < 0.0001 p < 0.0001
Bewick et al. 1999 (26)	57 MBC	High dose chemotherapy	TTP median 7.3 vs 14.3 mo OS median 17.7 vs 28.0 mo	p = 0.004
Bewick et al. 2001 (27)	46 MBC	High dose chemotherapy	TTP median 8.6 vs 13.0 mo	p = 0.009
Classen et al. 2002 (33)		2nd line chemo or hormonal	TTP median 13.9 vs 29.6 mo	p = 0.009 p = 0.002
Fehm et al. 1997 (38)	211 MBC	Anthracyclines and CMF	DFS median 11.0 vs 49.0 mo 5 vear OS 41% vs 61%	p = 0.01 p = 0.01
Fehm et al. 1998 (39)	80 MBC	CMF, NC or FNC	DFS less when elevated	p = 0.01
Fehm et al. 2002 (40)	52 PBC	Chemo or hormonal therapy	DFS 22 vs 27 mo	p = 0.04
	and 52 MBC			
Harris et al. 2001 (43)	425 MBC	High dose chemotherapy	DFS less when elevated OS less when elevated	p = 0.13 p = 0.0096
		AFM	DFS less when elevated	p = 0.061
Hayes et al. 2001 (44)	103 MBC	Chemo and hormonal therapy	OS median 16.4 vs 22.7 mo	p = 0.002 p = 0.002
Kandl et al. 1994 (48) Leitzel et al. 1995 (76)	79 MBC 300 MBC	Chemo and hormonal therapy Megestrol or fadrozole; 2nd line	OS median 21 vs 64 mo OS median 15.0 vs 28.0 mo	p = 0.030 p = 0.03 p < 0.0001

Lipton et al. 2002 (54)	719 MBC	Megestrol or fadrozole; 2nd line	TTP 90 vs 180 d TTF median 93 vs 175 d OS median 17.2 vs 29.6 mo	p < 0.0001 p < 0.0001 p < 0.0001 p < 0.0001
Lipton et al. 2003 (55) Mehta et al. 1998 (78)	562 MBC 79 PBC	Letrozole or tamoxifen CMF or CMFVP	TTP median 5.7 vs 9.4 mo DFS less when elevated	p = 0.0001 p = 0.045
Molina et al. 1996 (57)	412 PBC	dna	OS less when elevated DFS less when elevated	p < 0.0001 p < 0.0001
Narita et al. 1994 (61)	81 PBC	dna	OS less when elevated DFS less when elevated	p < 0.05
Nugent et al. 1992 (62)	161 PBC and	dna	OS less when elevated DFS less when elevated	p < 0.001 p < 0.001 p < 0.001
Willsher et al. 1996 (23)	6 MBC 81 PBC and	Tamoxifen	Stage I, II: OS median 27 vs >84 mo DFS median 11 vs >84 mo	p = 0.002 p = 0.002
Wu et al. 1999 (72)	38 MBC 226 PBC	Chemo and tamoxifen	Stage III: OS 24 vs 64 mo Stage IV: OS 18 vs 31 mo OS less when elevated	p = 0.04 p = 0.27 p = 0.0004
Yamauchi et al. 1997 (73)	94 MBC	Droloxifene 1st line	TTP shorter when elevated OS less when elevated	p = 0.001 p = 0.058
DNA, Data not available; N	1BC, metastatic	DNA, Data not available; MBC, metastatic breast cancer; PBC, primary breast cancer.	ar.	

5. SERUM HER2/neu AS A PREDICTIVE INDICATOR

5.1. Hormone Therapy

In the early 1990's Wright et al. (75), using IHC, reported that patients with metastatic breast cancer who demonstrated HER2/neu overexpression and estrogen receptor (ER) positivity, had a response rate of only 20% to first-line hormone therapy whereas 48% of the metastatic breast cancer patients who were ER positive but had a normal IHC expression of HER2/neu responded to the first-line hormone therapy.

Since that report there have been several publications concerning HER2/neu ECD levels and the response rate of metastatic breast cancer patients to first- and second-line hormone therapies. Table 2 lists several published studies representing 1778 metastatic breast cancer patients and 119 patients with primary breast cancer, and their response rates to hormone therapy with respect to serum HER2/neu ECD levels.

In the 2002 report by Lipton et al. (54), it was shown that MBC patients (n = 711) who were treated with second-line hormonal therapy (either the progestin megestrol acetate, or the aromatase inhibitor fadrozole) and who had a pretreatment serum HER2/neu level above the normal cutoff of 15 ng/mL were less likely to respond to the therapy (20.7%) compared to the 40.9% response of metastatic breast cancer patients who had serum HER2/neu levels below the normal cutoff of 15 ng/mL. The response rates reported here were nearly identical to those reported by the Wright et al. IHC studies mentioned earlier. The Lipton et al. studies also showed that patients with elevated pretreatment HER2/neu ECD levels had a shorter duration of response, a shorter TTP (illustrated in Fig. 4), and a shorter overall survival than ER-positive patients who had pretreatment serum ECD levels below the 15 ng/mL cutoff. In a recent report of first-line hormonal therapy, Lipton et al. (55) also demonstrated that metastatic breast cancer patients (n = 562) treated with either an aromatase inhibitor (letrozole) or an antiestrogen (tamoxifen) and who had elevated pretreatment serum HER2/neu levels, had a shorter time to progression, a shorter time to treatment failure, a decreased objective response rate (complete response plus partial response) and a decreased clinical benefit rate (complete response plus partial response plus stable disease greater than 24 wk) than similar patients who had serum HER2/neu levels below the normal cutoff of 15 ng/mL. Thus both forms of hormonal therapy were less effective if the pretreatment ECD levels were above the 15 ng/mL cutoff value. This study also showed that patients with normal serum HER2/neu ECD levels responded better to letrozole than to tamoxifen. However, there was no significant difference in response rate to letrozole or tamoxifen if the serum HER2/neu level was elevated. Therefore the superiority of letrozole was the greatest in those patients with normal serum HER2/neu levels.

Authors (ref.)	Patients	Therapy	Predictive indices (elevated vs nonelevated)	Significance
Hayes et al. 2001 (44)	103 MBC	Megestrol or fadrozole	RR 28% vs 37%	p = 0.041
Leitzel et al. 1995 (76)	300 MBC	Megestrol or fadrozole	RR 20.7% vs 40.9%	p = 0.004
			DR 11.6 vs 15.5 mo	p < 0.0001
Lipton et al. 2002 (54)	719 MBC	Megestrol, letrozole or fadrozole	RR 23% vs 45%	p < 0.0001
Lipton et al. 2003 (55)	562 MBC	Letrozole or tamoxifen	RR 15% vs 32%	p < 0.0001
			DR 18.5 vs 25.3 mo	p = 0.014
Willsher et al. $1999(23)$	119 PBC	Tamoxifen	RR 70% vs 70%	p = 0.71
Yamauchi et al. 1997 (73)	94 MBC	Droloxifene	RR 9% vs 56%	p = 0.00001
DR. Duration of response: RR. response rate.	S. response rate.			

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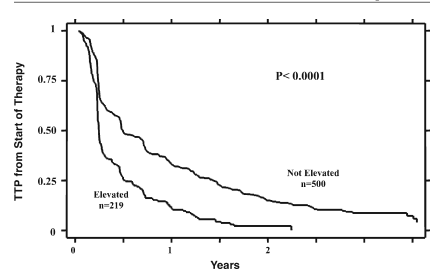


Fig. 4. Kaplan–Meier plot of TTP as a function of pretreatment serum HER2/neu level. These data are derived from three studies of patients with metastatic disease treated with second-line hormone therapy (two separate studies of fadrozole vs megestrol acetate; one study of letrozole vs megestrol acetate). HER2/neu not elevated, <15 ng/mL (n = 500) HER2/neu elevated, >15 ng/mL (n = 219).

5.2. Chemotherapy

Early studies by Gusterson et al. (79) and Muss et al. (80) reported that overexpression of HER2/neu correlated with a decreased responsiveness to combination therapy of cyclophosphamide/methotrexate 5-fluoro-uracil (CMF).

In 1995, Tsai et al. (81) demonstrated that an increased expression of p185neu resulted in enhanced chemoresistance following transfection of HER2/neu into non-small–cell lung cancer (NSCLC) cell lines established from untreated patients. In a follow-up 1996 report, the same investigators examined a panel of 20 NSCLC cell lines for HER2/neu ECD levels in the cell culture supernatant and then correlated the ECD levels with chemoresistance or chemosensitivity (82). Various cytotoxic drugs were used such as doxorubicin, cisplatin, etoposide, and malphalan. The results showed that high HER2/neu ECD levels were relatively chemosensitive. In this report, multivariate analysis revealed that the level of p185 neu was the only predictor for chemoresistance to doxorubicin and etoposide. In a 1997 report by Fehm et al., they showed that in node-positive breast cancer patients, those with elevated HER2/neu ECD levels had a worse outcome than patients

with nonelevated HER2/neu levels when treated with adjuvant CMF or cyclophosphamide–Novantrone–5-fluorouracil (38).

In 1997 Pegram et al. also reported that HER2/neu overexpression altered chemotherapeutic drug sensitivity in a variety of human breast and ovarian cancer cells (78). The next step in the analysis of the predictive role of HER2/neu expression and response to various regimens of chemotherapy was to examine published data concerning chemotherapy responses and HER2/neu ECD levels. These data are summarized in Table 3. We reviewed 12 publications, which contained data on 131 primary breast cancer patients and 1228 patients with metastatic disease.

In the report by Colomer et al. (34), 58 MBC patients were studied and it was shown that the probability of obtaining a complete response to a paclitaxel–doxorubicin chemotherapy regimen was significantly lower in patients with elevated HER2/neu ECD levels compared to patients with nonelevated levels. In addition, the duration of clinical response was significantly shorter in patients with elevated HER2/neu ECD levels compared with the cases with nonelevated levels. For instance the duration of response was only 7.5 mo for patients with ECD elevations compared to 11 mo for patients with normal ECD levels. Overall, elevated levels of the ECD correlated with reduced efficacy of a paclitaxel–doxorubicin chemotherapy combination.

In a report by Mehta et al. (83), it was concluded that breast cancer patients with more than three positive lymph nodes would benefit from determining the prechemotherapy levels of HER2/neu ECD and this could serve as an important marker to predict the response of breast cancer patients to chemotherapy. Postchemotherapy c-*erb*-B2 levels were also a prognostic indicator for disease-free survival in patients who received chemotherapy. Both the Mehta et al. (83) and Fehm et al. reports (40) concluded that patients with elevated HER2/neu ECD levels had a lower response to first-line therapy than breast cancer patients with normal serum HER2/neu ECD levels.

Nunes and Harris (11) as well as Kaptain et al. (10) reviewed the role of HER2/neu ECD in relation to alkylating-containing regimens, anthracycline-containing regimens, and taxanes. Overall they concluded that HER2/neu–positive tumors are relatively resistant to CMF-containing regimens but had an increased sensitivity to anthracycline-containing regimens. The clinical value of HER2/neu ECD testing and response to various chemotherapeutic regimens continues to be an area of research investigation. Therefore, the availability of an FDA-cleared serum HER2/neu ECD test will allow standardized studies to be conducted for the comparison of data between laboratories.

Yu et al. (84) showed that HER2/neu-transfected cells are more resistant to paclitaxel. However, the effect of taxanes in xenograft models in conjunc-

Summary of D	Data on HER2	/neu as a Predictor of Response to Cl	Summary of Data on HER2/neu as a Predictor of Response to Chemotherapy in Patients With Breast Cancer	er
Authors (ref.)	Patients	Therapy	Predictive indices (elevated vs nonelevated)	Significance
Bewick et al. 1999 (26) Classen et al. 2002 (33)	<i>57</i> MBC 64 MBC	HDCT and ABSC Tamoxifen, adriamycin+	DFS 7.5 vs 14.3 mo RR 11.5% vs 84.2%	p = 0.004 p < 0.001
Colomer et al. 2000 (34)	58 MBC	cyclophospan or CMF Paclitaxel and doxorubicin	CR 0% vs 26% No recronce 37% vc 73%	1 CU U
Fehm et al. 1997 (38)	211 MBC	NC or FNC, CMF	DFS 11.0 vs 49.0 mo	p = 0.021 p = 0.01
Fehm et al. 1998 (39)	80 MBC	CMF, NC or FNC	RR 29% vs 59%	
Fehm et al. 2002 (40)	52 PBC	Chemo and hormonal therapy	80% levels corresponded with response	
	and 52 MRC			
Harris et al. 2001 (43)	425 MBC	Doxorubicin. 5FU. methotrexate	Lower response when elevated	n = 0.061
Hayes et al. 2001 (44)	139 MBC	Anthracyclines	RR 29% vs 38%	p = 0.26
Kandl et al. 1994 (48)	74 MBC	No details	RR 50% vs 50%	NS
Luftner et al. 1999 (56)	35 MBC	Dose intensified paclitaxel	Higher response (69%) when elevated	
1000 11		monotherapy	1	
Menia et al. 1996 (70) Revillon et al. 1996 (63)	33 MBC	CIMIF, CIMIF V F Vinorelbine	Lower response when elevated RR 90% vs 61%	p = 0.04 0.5 < n < 0.9
CD Comulate recuonce: D	DFC disease fre	OD Comulata recnonce: DEC diceases free curvivel. DD recnonce rete		

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CR, Complete response; DFS, disease-free survival; RR, response rate.

tion with Herceptin appeared to reverse the resistance to both paclitaxel and docetaxel (86), which is especially interesting because some of the best current response rates are seen when Herceptin is combined with docetaxel or paclitaxel (86,87).

5.3. Herceptin-Based Therapy

In a report in 2002, Esteva et al. analyzed the response rate of 30 MBC patients (all shown to be HER2/neu-positive by tissue testing) treated with docetaxel and Herceptin who were separated into responders with high or low ECD levels (86). The circulating HER2/neu ECD levels were measured at baseline and at the time of response evaluation for all 30 patients. The median level at baseline was 41.9 ng/mL (range, 7.1 ng/mL to 666.5 ng/mL). Twenty-one patients (70%) had elevated HER2/neu levels at baseline. The patients with high ECD baseline levels had the highest response rate to Herceptin-based therapy (76%) whereas only 33% of those with low ECD levels responded. In comparison, 67% of the FISH-positive patients responded. The overall response rate achieved was 63%; however, when patients with minor responses and stable disease were considered, 83% of the patients obtained some clinical benefit from the combination of weekly docetaxel and Herceptin. Esteva et al. also reported that serial changes in serum HER2/neu ECD levels correlated very well with clinical response to weekly docetaxel and Herceptin therapy. The authors concluded that additional research is warranted to determine the value of serum HER2/neu ECD testing in selecting and monitoring patients undergoing Herceptin-based therapy.

In a retrospective study designed to determine whether HER2/neu levels could predict outcomes of Herceptin-based therapy, Hoopman et al. (87) examined plasma samples from 20 MBC patients. HER2/neu ECD levels were measured in samples collected at the beginning of Herceptin-based therapy and at the time of first diagnostic imaging. HER2/neu levels were evaluated in relation to the clinical course of each patient and using imaging results, and then the data were analyzed with respect to prediction of the patient response to therapy defined as the time to progression. The data presented showed that the change in plasma HER2/neu levels were predictive of clinical course in 16 of 20 patients (80%) and that all therapy responders had normal ECD levels at the time of first diagnostic imaging. The patients with permanently elevated or increasing serum HER2/neu levels displayed a poor clinical outcome in 6 of the 10 cases. Based on these results the authors concluded that plasma HER2/neu levels might be an early predictor of response to Herceptin-based therapies.

Finally, in a report by Dnistrian et al. (37) metastatic breast cancer patients were stratified according to pretreatment serum HER2/neu levels to determine whether ECD levels could predict the response to Herceptin-based

therapies. Of 18 patients treated with Herceptin and Taxol who had pretreatment ECD levels less than the cutoff (15 ng/mL), 6 responded favorably to Herceptin-based therapy. In contrast, 30 of the 36 patients (83%) who had elevated pretreatment serum HER-2/neu levels (mean 344 ng/mL), had a favorable response. Furthermore, in all patients with abnormally high pretreatment serum HER2/neu levels, the ECD level decreased significantly with disease regression and in most cases it returned to normal.

The observation that elevated serum HER2/neu levels may be predictive of response to Herceptin-based therapy (summarized in Table 4) is interesting in light of the publications demonstrating that elevated pretreatment HER2/neu ECD levels predict poor response to hormone therapy and some regimens of chemotherapy. It is not entirely unexpected, however, that high levels can predict response to Herceptin because the HER2/neu receptor provides the binding site for Herceptin which is necessary for its action. The shed ECD may therefore provide an indication of the amount of receptor that is available for binding, as well as the activity of the malignancy. Additional studies will need to be done to clarify this possibility.

6. MONITORING HER2/neu ECD LEVELS IN MBC PATIENTS TREATED WITH HORMONE OR CHEMOTHERAPY

In this section (summarized in Table 5), we reviewed 16 references (14 related to MBC and 4 related to PBC) representing 1148 breast cancer patients. The 1148 patients were divided into 499 MBC patients and 649 primary breast cancer patients. In these studies serial or longitudinal changes in serum HER2/neu ECD values were compared to the clinical course of disease in women with MBC. The clinical course of a patient's disease (defined as progression, response or stable disease) was determined by monitoring results of clinical tests such as X-ray films or CT scans. Patients were then classified as to whether changes in serum HER2/neu levels did or did not correspond to the clinical course of disease. For example, it was determined whether the serum value increased with progression or decreased with response to therapy to determine correspondence.

In a report by Cook et al. (35), longitudinal monitoring was performed on 103 stage IV breast cancer patients who were being treated with various regimens of hormonal or chemotherapy, to determine whether serum HER2/neu changes correlated with changes in the clinical course of disease. Thirty-eight (36.9%) of the 103 stage IV patients had an elevated ECD, 33 of whom showed longitudinal HER2/neu values which paralleled the clinical course of disease, thus giving an overall sensitivity of 86.8%.

Schwartz et al. (65) reported serum HER2/neu levels in patients with MBC who were receiving a variety of conventional therapeutic regimens.

Summary of Data on Serum I	HEKZ/neu as a I	redictor of Kesponse to Hercel	Summary of Data on Serum HEK2/neu as a Predictor of Response to Herceptin-Based 1 herapies in Patients With Metastatic Breast Cancer	breast Cancer
Authors (ref.)	Patients	Therapy	Prognostic indices	Significance
Dnistrian et al. 2003 (<i>37</i>) Esteva et al. 2002 (<i>86</i>) Hoopmann et al. 2003 (<i>87</i>) Schoendorf et al. 2002 (<i>64</i>)	54 MBC 30 MBC 20 MBC 23 MBC	Herceptin ± paclitaxel Doxetaxel and herceptin Various chemotherapies and herceptin Various chemotherapies and herceptin	RR 86% in elevated, 61% in nonelevated RR 76% in elevated, 33% in nonelevated Predicted clinical course in 80% of patients Corresponded with clinical course in 74% of clinical events	p < 0.001 p = 0.04
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RR, Response rate.

Summar	y of Data on	HER2/neu for Monitoring Ther	Summary of Data on HER2/neu for Monitoring Therapy and Early Detection of Metastases	
Authors (ref.)	Patients	Therapy	Monitoring observation	Significance
Cook et al. 2001 (35)	38 MBC	No details given	86.8% concordance between changes and clinical course	
Cheung et al. 2000 (<i>32</i>)	30 MBC	Docetaxel/Doxorubicin based	100% concordance with course in all with elevated levels Earlier detection of progression;	
Dnistrian et al. $2003 (37)$	54 MBC	Herceptin ± paclitaxel	87.5% concordance between changes	
Isola et al. 1994 (47)	8 MBC	No details given	100% concordance between changes and clinical course	
Fehm et al. 2002 (40)	52 PBC and 52 MBC	Chemo or hormonal therapy	27% and 50% of patients had elevated HER2/neu 6 and 3 mo	
Hoopmann et al. 2003 (<i>87</i>)	20 MBC	Various chemotherapies and herceptin	Predicted clinical course in 80% of patients Level increased in 35% of patients at time	
Kath et al. 1993 (50)	8 MBC	Various chemo and hormonal therapies	or detection of interastases by intraging 100% concordance with clinical course	
Luftner et al. 1999 (56)	35 MBC	Dose intensified Paclitaxel	Levels follow clinical course	

Table 5

<i>p</i> = 0.007			
Elevated levels associated with relapse, lead time 6–9 mo	Lead time to detection of relapse 2-9 into Lead time to relapse 4.5 \pm 2.5 mo Lead time to relapse 4.8 \pm 2.4 mo Levels followed clinical course Levels paralleled course in 74% of events	Levels reflect clinical course	Indicator of relapse sens 58.3% spec 85.2% 58% of patients showed concordance with clinical course
No details given	No details given No details given No details given Various chemo and hormonal therapies	and herceptin Various chemo and hormonal therapies and hercentin	Tamoxifen, CAF Fadrozole, megestrol acetate
18 MBC	200 PBC 250 PBC 4 MBC 23 MBC	147 PBC and 138 MBC	3 MBC 48 MBC
Mansour et al. 1997 (88)	Molina et al. 1996 (57) Molina et al. 1999 (60) Narita et al. 1992 (61) Schoendorf et al. 2002 (64)	Schwartz et al. 2000 (65)	Sugano et al. 2000 (68) Volas et al. 1996 (22)

All patients had a pretherapy serum specimen and four posttherapy specimens. Longitudinal testing of the serum HER2/neu levels showed clearly that women who expressed HER2/neu in their tissue had elevated serum levels and that changes in the ECD levels reflected the clinical course of the patients' disease. This report also investigated whether there were serial changes in normal individuals over a several month period. In the study, six specimens were drawn monthly from seven premenopausal and eight postmenopausal women and tested for serum HER2/neu levels. These studies demonstrated that the serial HER2/neu values were consistent for a given person over a several month period. In one representative example, the serum HER2/neu values over several months were 6.2 ng/mL, 8.3 ng/mL, 7.7 ng/mL, 8.3 ng/mL, 8.1 ng/mL, and 8.1 ng/mL. Similar data was demonstrated with the remaining normal individuals, and was therefore a good basis for comparative monitoring studies in MBC patients.

Cheung et al. (32) examined serial changes in serum HER2/neu levels in 30 MBC patients from 2 multicentered trials in which patients received either docetaxel-based therapy or doxorubicin-based therapy. The authors concluded that among the patients with positive tissue staining, sequential changes in serum completely paralleled the initial response to therapy.

In a report by Lueftner et al. (56), serum samples were taken weekly from 35 patients to monitor changes in the serum HER2/neu ECD levels and to correlate those changes to the clinical course of disease. The MBC patients received dose-intense paclitaxel treatment. In this study the overall response rate was 36% but the response rate among the HER2/neu-positive patients was 62%, showing a high sensitivity of the HER2/neu-positive patients to dose-intense paclitaxel treatment. In all responders, the HER2/neu level decreased below the detection limit either before the clinical diagnosis of response or by the end of the next cycle. However, in this cohort of patients normalization of the HER2/neu levels also occurred in patients that were stable or had progressive disease. The authors speculate that this could be explained by the fact that the chemotherapy is effective against the HER2/neu-positive tumor cells but not against the HER2/neu-negative tumor cells that could be responsible for the progressive disease. It was also pointed out by the authors that chemotherapy may alter the mechanisms by which the ECD is shed, but no data was presented to support this hypothesis.

In summary, numerous reports of patients receiving hormone or chemotherapy showed that longitudinal changes in serum HER2/neu levels paralleled the clinical course of a patient's disease. Overall, several studies showed that increases in serum HER2/neu levels were reflective of progressive breast cancer while decreasing serum HER2/neu levels were reflective of response to therapy or a prolonged lack of disease progression.

7. MONITORING METASTATIC BREAST CANCER PATIENTS TREATED WITH HERCEPTIN-BASED THERAPIES

In studies presented in Table 5, women with MBC being treated with Herceptin and various chemotherapies (n = 110) were monitored for changes in HER2/neu ECD levels. HER2/neu ECD levels were determined prior to treatment and then serially thereafter. Previous studies reported by Payne et al. have shown that Herceptin does not interfere with measuring ECD levels in the HER2/neu assay used in all three of these studies described below (74). In an initial report by Schwartz et al., in 2000, it was suggested that changes in serum HER2/neu during Herceptin-based therapy might parallel the clinical course of disease; however, there were too few patients in the report to make a valid conclusion (65).

In a 2002 report by Esteva et al. (86), 30 MBC patients treated with docetaxel (Taxotere) and Herceptin were monitored for changes in serum HER2/neu to see if serial changes would reflect the clinical course of disease. Studies showed that ECD levels decreased in 14 of 16 (87%) of responding patients. As seen in studies of patients receiving conventional therapies, serial changes in serum HER2/neu levels did parallel the clinical course of disease after Herceptin treatment.

In a report by Schoendorf et al. (64), 23 MBC patients treated with Herceptin and chemotherapy (for a median time of 13 mo, with a range of 4–22) were monitored serially for the change in serum HER2/neu ECD. The changes in HER2/neu ECD level were then evaluated in conjunction with response or lack of response to therapy. In the group of patients with elevated levels (12 of 19), 35 events of either response or progression were documented. The serial changes in serum HER2/neu correlated with remission or disease progression in 74% of the patients. The correlation between serial ECD changes and clinical changes was increased when the analysis was focused on MBC patients with visceral metastasis. This group of clinical investigators concluded that serial changes in plasma HER2/neu ECD levels did parallel changes in the clinical course of disease.

In a 2003 report (89) a group of 57 MBC patients treated over a 2-year period with Herceptin and Taxol were monitored for changes in serum HER2/neu and the data was correlated with clinical changes. The studies clearly showed that serum HER2/neu ECD changes paralleled the clinical course of disease. The women with serially decreasing ECD levels responded to Herceptin-based therapy while women with progressing breast cancer that did not respond to the combined therapy had serially increasing ECD levels. These data were consistent with the observations made in MBC patients who were treated with conventional hormone and chemotherapies.

In summary, these four publications collectively studied a small number of women with MBC who received Herceptin-based therapies and who had their serum HER2/neu levels monitored for up to 2 yr. All four publications showed that serial changes in the ECD levels reflected the clinical course of disease. Women who had serially decreasing ECD levels responded to Herceptin-based therapy whereas women with progressive breast cancer had serially increasing ECD levels.

8. SERUM HER2/neu LEVELS AND DETECTION OF EARLY RECURRENCE

In the 1996 report by Molina et al. (58), they evaluated the utility of measuring HER2/neu, carcinoembryonic antigen (CEA), and CA 15-3 in the early diagnosis of recurrence. Serial serum measurements were performed in 200 primary breast cancer patients (no evidence of residual disease) followed for 1–4 yr with a median of 2.2 yr. Of the 89 patients who developed metastasis, 28% had a serum HER2/neu above the cutoff. 30% had a CEA that was above the normal cutoff and 47% demonstrated elevated levels of CA 15-3. The lead time prior to diagnosis was $4.5 \pm$ 2.4 mo for HER2/neu, CEA was 4.9 ± 2.4 mo and CA 15-3 was $4.8 \pm$ 2.4 mo. However, sensitivity was clearly related to the site of recurrence, with the lowest sensitivity found in locoregional relapse and the highest in patients with visceral metastasis. When patients with locoregional relapse were excluded, the sensitivity for HER2/neu improved to 31% and the overall sensitivity of early detection with all three markers combined was 76%. This data and others are summarized in Table 5. The increased sensitivity observation was also supported by Watanabe (21) and Schwartz (65), however, Eskilinen only reported limited value in measuring serum CEA. CA 15-3, and HER2/neu in conjunction with other cancer tests (90).

In the study by Dnistrian et al. (*37*) HER2/neu, CA 15-3 and CEA were all measured serially from baseline to investigate the changes in 54 metastatic breast cancer patients undergoing Herceptin and Taxol therapy. When the individual data was combined the concordance with monitoring for response was 76%, similar to Molina et al. (*58*), but individually the concordance was 67% (only 31% in the Molina report) for HER2/neu, 54% for CA 15-3 and 43% for CEA.

In a report by Ali et al. (24), a study was done to measure CA 15-3 (a surrogate marker for disease burden) with serum HER2/neu in 566 ER/PR-positive metastatic breast cancer patients. These patients were treated with second-line hormone therapy, megestrol acetate, or an aromatase inhibitor, Fadrozole. Overall, 30% of the patients had an elevated HER2/neu ECD level (n = 168) and 60% had an increased CA 15-3; however, there was only a weak correlation between the two. Similar to a previous report, the clinical

benefit (complete response plus partial response plus stable disease greater than 24 wk) of endocrine therapy was significantly lower in patients with elevated HER2/neu ECD levels. The investigators concluded that HER2/ neu was a significant independent predictive and prognostic factor in hormone receptor-positive MBC patients even when adjusted for tumor burden as measured by CA 15-3. The combination of an elevated HER2/neu and CA15-3 predicted a worse prognosis for MBC patients than did an elevated CA 15-3 alone.

The Colomer report (34) suggests that a panel of tumor tests such as CEA, CA 15-3 and HER2/neu could be used to monitor patients postoperatively to increase the sensitivity of detecting early recurrence. The value of early detection will increase with the introduction of a variety of targeted therapy used as either monotherapy or in conjunction with hormonal or chemotherapies.

9. CONCLUSIONS

This chapter has shown that the prevalence of an elevated serum HER2/neu ECD is highly variable in breast cancer, which probably reflects the differing time of sampling in relation to the evolution and aggressiveness of the disease. As might be expected, there is a higher prevalence of elevated levels in the metastatic phase of the disease, conforming with the view that increased HER2/neu amplification is associated with a more aggressive form of the disease. There is strong data to show that an elevated serum HER2/neu ECD is an indicator of poor prognosis and furthermore is a predictor of poor response to therapy using chemotherapeutic and hormonal treatment regimens. Conversely an elevated serum HER2/neu ECD level is a predictor of improved response to Herceptin-based therapy, as it reflects the presence of an increased population of target molecules for the drug to bind to the malignant cell.

This chapter has also demonstrated an interesting observation regarding the potential use of serum HER2/neu ECD measurement as a tool for detecting the development of metastatic disease, ahead of conventional indicators, as well as monitoring of response to a variety of therapies administered in the metastatic phase of breast cancer. The review has also clearly established the need for more studies to clarify the clinical value of HER2/neu ECD testing in breast cancer patients. The review has also established the relationship between serum HER2/neu ECD levels and the presentation of breast cancer. It has identified several potential roles for the HER2/neu oncoprotein in clinical decision-making with respect to the diagnosis and management of breast cancer. In turn this could lead to improvement in clinical outcome. Prospective studies are also required in which the use of the HER2/neu oncoprotein test is used as a decision-making tool to demonstrate improved clinical outcomes.

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14

Circulating Vascular Endothelial Growth Factor

Methods, Prognostic Significance, and Potential Application for Antiangiogenic Therapy

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CONTENTS

INTRODUCTION ROLE OF ANGIOGENESIS IN TUMOR GROWTH AND PROGRESSION DETERMINATION OF TISSUE ANGIOGENESIS AS A CANCER BIOMARKER CIRCULATING BIOMARKERS OF ANGIOGENESIS: PROMISES AND PITFALLS CONCLUSIONS REFERENCES

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SUMMARY

Despite significant advances in early detection and treatment, breast cancer still remains the major cause of cancer-related death in women. Many studies suggest a relationship between angiogenesis and breast cancer prognosis. Angiogenesis is the complex process leading to the formation of new blood vessels from pre-existing vascular network. The VEGF is the most active growth factor involved in angiogenesis; more specifically, raised intratumoral VEGF concentrations have been shown to correlate with tumor aggressiveness. VEGF is therefore a promising target for new therapies, but it is still unclear in which blood matrix the determination of VEGF is more accurate as a cancer biomarker and which matrix provides the optimal clinical information. Circulating levels of VEGF have been measured by several investigators who reported conflicting results. However, these studies are not comparable with each other due to a lack of standardization of the pre-analytical phase. The chapter presents the main studies concerning anti-VEGF therapies; several studies evaluated the safety profile and activity of the combination of standard chemotherapy with new antiangiogenic agents. However, to date only a few definitive results on the effect of angiogenesis blood markers have been reported. Determination of circulating VEGF still remains an experimental procedure with no evident application for routine clinical decisions. Data from retrospective studies, however, suggest that VEGF levels may predict clinical outcome of breast cancer.

Key Words: Vascular endothelial growth factor; breast cancer; angiogenesis; therapy; prognosis.

1. INTRODUCTION

Angiogenesis, or neovascularization, is the complex process leading to the formation of new blood vessels from the preexisting vascular network of the tissue. It is necessary for new organ development and differentiation during embryogenesis, wound healing, and reproductive functions in adults (1). Angiogenesis is also involved in pathogenesis of certain chronic diseases, such as rheumatoid arthritis, age-related macular degeneration, proliferative retinopathies, psoriasis, as well as malignant tumors. Angiogenesis is a complex process that, during physiologic tissue growth and repair, is closely regulated by pro- and anti-angiogenic growth factors (Table 1). In cancer, the net balance of pro- and antifactors is altered vs down-regulation of angiogenic inhibitors or up-regulation of angiogenic activators, providing tumor progression and metastasis. Neovascularization occurs in a series of complex and interrelated steps. First, tumor or certain stromal cells (e.g., macrophages, mast cells, and fibroblasts) release endothelial cell

Proangiogenic factors	Naturally occurring inhibitors
Angiogenin	Angiostatin (plasminogen fragment)
Angiopoietin-1	Antiangiogenic antithrombin III
Del-1	Cartilage-derived inhibitor (CDI)
Fibroblast growth factor, acidic (aFGF)	CD 59 complement fragment
Fibroblast growth factor, basic (bFGF)	Endostatin (collagen XVIII fragment)
Follistatin	Fibronectin fragment
Granulocyte colony-stimulating factor (G-CSF)	Gro-β
Hepatocyte growth factor (HGF)	Heparinases
Interleukin-8 (IL-8)	Heparin hexasaccharide fragment
Leptin	Human chorionic gonadotropin (hCG)
Midkine	Interferon- α , β , γ
Placental-growth factor (PiGF)	Interferon inducible protein (IP-10)
Platelet-derived endothelial cell growth factor	Interleukin-12 (IL-12)
Platelet-derived endothelial cell growth factor-BB	Kringle 5 (plasminogen fragment)
Pleiotrophin (PTN)	Tissue inhibitors
1 ()	of metalloproteinases
Proliferin	2-Methoxyestradiol
Transforming growth factor- α (TGF- α)	Placental ribonuclease inhibitor
Transforming growth factor- β (TGF- β)	Plasminogen activator inhibitor
Tumor necrosis factor- α (TNF- α)	Platelet factor 4 (PF4)
Vascular endothelial growth factor	Prolactin 16-kDa fragment
(VEGF)	Retinoids
	Tetrahydrocortisol-S
	Thrombospondin-1
	TGF-β
	Vasculostatin
	Vasostatin

Table 1 Proangiogenic and Antiangiogenic Factors

growth factors, such as vascular endothelial growth factor (VEGF), fibroblast growth factor (FGF), or platelet-derived growth factor (PDGF) into the surrounding tissue. These factors are secreted in response to proteins overexpressed in the microenvironment for example, epidermal growth factor (EGF), insulin growth factor (IGF), FGF, interleukins, PDGF, or in response to several mechanisms as hypoxia, hypoglycemia, inflammation, and genetic alterations. The endothelial growth factors bind to, thereby activating, endothelial cells that form the walls of new blood vessels. Subsequently, activated endothelial cells stimulate proteolytic enzymes that break down the extracellular matrix, allowing endothelial cells to invade the matrix and migrate. The proliferation and migration of activated endothelial cells give origin to new capillary tubes. Endothelial cells differentiate and synthesize a new basement membrane (2). The maturation and differentiation of the new vessel is completed with the formation of the vascular lumen. Finally, the adhesion receptor integrins $\alpha v\beta$, present on the surface of activated endothelial cells, permit the linkage of the new vessels with the preexisting ones to generate the intratumoral vascular network. The angiogenic process is necessary for tumor growth, invasiveness, progression, and metastasis and its determination has multiple important clinical applications as a potentially useful prognostic indicator, or as predictive marker of response to antiangiogenic therapy.

2. ROLE OF ANGIOGENESIS IN TUMOR GROWTH AND PROGRESSION

Despite significant advances in early detection and treatment, breast cancer still remains the major cause of cancer-related death in women.

There are compelling data suggesting a relationship between angiogenesis and breast cancer prognosis. Preclinical studies in vivo have shown that while normal, healthy breast tissue has no angiogenic activity, all breast carcinoma samples have some degree of angiogenesis (3).

The growth of MCF-7 cells transfected to overexpress VEGF in vivo correlates with a significant increased number of microvessels, indicating the importance of neovascularization for tumor growth (4). The experimental findings are supported by clinical studies that have demonstrated the prognostic significance of angiogenesis in breast cancer, either in terms of metastatic potential (5), relapse-free survival (6), or long-term survival (7). Most of the studies have found that an increased level of angiogenesis, as measured by the assessment of microvessel density, is associated with a reduced survival, shorter relapse-free survival, and an increased risk of metastases. Indirect markers of angiogenesis, such as microvessel density, could potentially be used to select those patients with a poor prognosis for whom alternative therapeutic approaches may be necessary.

The "angiogenic switch" is characterized by oncogene-driven tumor expression of pro-angiogenic proteins, such as VEGF, bFGF, interleukin-8, transforming growth factor- β , PDGF, and others (8–10). It has been shown that other mediators of neovascularization such as interleukins, oncogenes, and tumor growth factors may produce their effects by altering the expression of VEGF, suggesting that the VEGF pathway plays a central role in in vivo angiogenesis (11).

Among the endothelial cell growth factors, VEGF is the most active, specific, and potent mitogen for vascular endothelium (12). VEGF is a promising target, as it has been shown to be a potent promoter of endothelial cell proliferation (13) and chemotaxis (14), and it also increases vascular permeability (15). VEGF-A is a dimeric 34–42-kDa glycosylated basic protein, encoded in five isoforms; the two larger isoforms (VEGF 189 and VEGF 209) are cell associated, while the two smaller ones (VEGF 121 and VEGF 165) are secreted as soluble molecules. VEGF activation requires the phosphorylation of specific transmembrane tyrosine kinase receptors, which are preferentially expressed on vascular endothelial cells.

Three specific VEGF related tyrosine-kinase receptors have been identified: VEGF-R1 (flt-1 or fms-like-tyrosine kinase), VEGF-R2 (flk-1/KDR), located on vascular endothelial cells (Ecs) and VEGF-R3 (flt-4), located on lymphatic vessels (*16–18*). Recently, two isoform-specific nontyrosine kinase receptors, neurophilins 1 and 2, involved in neural cells guidance, have also been reported to bind VEGF (*19*). Neurophilin 1 binds VEGF-165, but not VEGF 121, and neurophilin 2 is correlated to VEGFR-1 (*20*). Activated VEGFR phosphorylates several signaling cascade proteins, including phospholipase C, phosphoinositol-3 kinase, and ras GTPase activating proteins (*21*). This cascade of cellular events leads to proliferation, migration, and differentiation of activated endothelial cells (*22*).

The VEGF family includes at least sequenced isoforms, derived by alternative exone-splicing: VEGF 121 (A), 165 (B), 189 (C), 206 (D), and 145 (E). Three other members—PIGFs 1, 2, and 3—are also described as alternative spliced isoforms of the same gene (12,23). Recently, four independent research groups provided direct evidence that VEGF-C and VEGF-D are important regulators of lymph vessel growth in vivo, both being ligands for the VEGF-3 receptor (16,17). These studies provided the first experimental evidence that tumors are able to promote lymphangiogenesis, a phenomenon that has been neglected for a long time. By using a novel specific marker for lymphatic endothelium, the anti-LYVE-1 antibody (24), a significant correlation of lymphatic vessel immunostaining with overexpression of VEGF-C or VEGF-D was demonstrated. Indeed, tumor lymphangiogenesis was associated to lymph node metastasis. Skobe et al. (17) demonstrated the occurrence of intratumoral lymphangiogenesis within human breast cancer after orthotopic transplantation onto nude mice. The degree of lymphatic vessels density was associated with overexpression of VEGF-C and enhanced regional lymph node and lung metastasis. Moreover, another study (25) showed that VEGF-C overexpression is detectable only in human invasive breast cancers with histologically proven axillary lymph node metastasis. Makinen et al. (26) found in an experimental model that a soluble VEGFR-3 fusion protein inhibits the process of lymphatic vessel development, leads to regression of existing lymphatics in vivo, and reduces lymphedema in transgenic mice expressing soluble VEGFR-3.

Because the evaluation of a blood sample is easier in comparison to immunohistochemical assessments, or immunoassays that involve laborious tissue preparation procedures and it allows for serial measurements, many studies have been published on circulating VEGF, as a surrogate marker of angiogenesis.

VEGF gene expression is regulated by several mechanisms: hypoxia, glucose deprivation, soluble cytokines, oxidative and mechanical stresses, oncogenes, and tumor-suppressor genes mutations (27,28).

3. DETERMINATION OF TISSUE ANGIOGENESIS AS A CANCER BIOMARKER

Studies have confirmed the presence of VEGF gene overexpression in malignant tumors as compared to benign breast tissues. Raised intratumoral VEGF concentrations have also been shown to correlate with tumor aggressiveness. However, there are conflicting reports on the association of VEGF levels with disease-free survival (29–31).

The clinical significance of cytosolic VEGF levels was first tested by Gasparini et al. in two studies published in 1997. The first study (32) evaluated VEGF protein in 260 consecutive patients with node-negative disease not treated with adjuvant therapy, median follow-up of 72 mo. In both univariate and multivariate analysis for RFS and OS, VEGF retained a significant and independent prognostic value. The second study (33) was performed in the same cohort of cases, with prognostic evaluation extended also to other biological factors, such as: cathepsin D, p53 protein, and TP. More recently, Gasparini et al. (34) performed a study to evaluate the clinical significance of co-determination of VEGF and TP in series of nodepositive breast cancer patients treated with adjuvant therapy. Two series of patients were evaluated: the first group of patients included 137 patients treated with adjuvant chemotherapy (CMF iv schedule). The second group included 164 patients who received adjuvant hormone therapy (tamoxifen). In the first group of patients, the two angiogenic peptides were found to be significant and independent prognostic factors. In multivariate analysis on RFS only VEGF and the number of involved lymph nodes retained a significant and independent prognostic value. Both estrogen and progesterone receptors were significant prognostic indicators for both RFS and OS in univariate analysis, but they lost their significance in the multivariate model inclusive of VEGF. No statistically significant associations were found between VEGF and the other prognostic factors examined (age, menopausal status, histologic tumor type, tumor size, and hormone receptors). The lack of association of VEGF with hormone receptors suggests that this angiogenic peptide is likely to stimulate the growth of human breast cancer independently of the hormone pathways via direct autocrine/or paracrine

stimulation on tumour cells or enhancing intratumoral vascular permeability, thus allowing more oxygen and nutrient to reach the tumor.

It has been recently hypothesized that overexpression of VEGF by estrogen-dependent breast cancer cells could produce an effect on breast cancer progression (acquisition of estrogen-independent growth) similar to the growth stimulation induced by estrogens. Li et al. (35) found that overexpression of the VEGF isoforms 121 and 165 by estrogen-dependent MCF-7 breast cells stimulated breast tumor formation in an estrogenindependent fashion in ovariectomized mice, in the absence of estrogen treatment. In addition, VEGF strongly stimulated neovascularization in MCF-7 tumors either in estrogen-treated or untreated mice, as well as enhanced estrogen-dependent tumor growth in estrogen-treated mice. These findings suggest that up-regulation of VEGF indirectly contributes to the acquisition of estrogen-independent cancer growth by stimulating tumor angiogenesis.

4. CIRCULATING BIOMARKERS OF ANGIOGENESIS: PROMISES AND PITFALLS

4.1. Methods of Determination: Factors Impacting on Clinical Value of the Biomarker

4.1.1. PREANALYTICAL REASONS FOR VARIABILITY

Circulating levels of VEGF have been measured by several investigators who reported conflicting results (Table 2). However, these studies are not comparable owing to a lack of standardization of the preanalytical phase (*36*). In fact, the utility of biomarkers in oncology is closely dependent to the information that comes from its measure. It is therefore necessary to define standardized operative procedures to be used in all the phases of determination (preanalytical, analytical, and postanalytical).

It is still unclear in which blood matrix the determination of VEGF is more accurate as a cancer biomarker and which matrix provides the optimal clinical information. Circulating VEGF has a multicompartmental origin because it is mainly produced by cancer cells and/or transported and released by platelets, lymphocytes, granulocytes, monocytes, and megakaryocytes. Consequently, the different compartments in which VEGF is measured such as serum, plasma, or whole blood, may give different information.

The presence of VEGF in the alpha granules of platelets (*37*) implies that serum VEGF levels not only reflect the circulating VEGF, but also the VEGF released from blood cells during the coagulation process. Normally, plasma of healthy people does not contain relevant quantities of VEGF.

As a consequence, any variation in sample handling may affect the blood cell activation and, thus, the release of VEGF into serum. Activated platelets release VEGF in a rapid discharge reaction, increasing the VEGF content up

Table 2 Summary of Methods for Sample Collection Reported in the Literature

			a. S	a. Serum						
		Temnerature	Clotting	8ı	Cen	Centrifugation	ı	Serum	VE in	VEGF levels found in health controls
Authors (ref.)	Collection tube	sample blood	Time	Temp.	G	Time	Temp.	storage	Cases	VEGF (pg/mL)
Yamamoto 1996 (49)	NR	NR	NR	NR	3000 rpm 10 min	10 min	4°C	-20°C	184	0–227.5 Min–max
Dirix 1997 (94)	Vacutainer system	RT	NR	NR	3000 rpm	10 min	NR	-80°C	NR	Mean = // NR
Salven 1997 (42)	Venoject blood	4°C	30 min	4°C	2000 g	10 min	4°C	-70°C	113	1-177
Verheul 1997 (38)	conection system (Terumo) NR	NR	20–30 min	RT	3000 rpm	NR	NR	-20°C	30	Median = 15 0-287
Banks 1998 (39)	Plastic tube Monovette	RT	NR	NR	2000g	10 min	NR	-70°C	~	Min-max 76-854 Min-may
Benoy 1998 (95)	(Sarstedt) NR	NR	NR	NR	NR	NR	NR	-80°C	9	30-403 Min-max
Heer 1998 (47)	NR	NR	NR	NR	NR	NR	NR	-80°C	14	Median = 153 83.1-327.7
Maloney 1998 (96)	Plastic tube	RT	30 min	22°C	750g	10 min	4°C	-70°C	10	1000 ± 63 230 ± 63 00000 ± 6D)
Webb 1998 (97)	(Decton Dickinson) Plain glass	RT	2 h	NR	NR	NR	NR	-80°C	34	Mean = 249.4
Agrawal 1999 (48)	(Beccoll DICKIIISOII) NR	NR	NR	NR	NR	NR	NR	-80°C	14	1300–7000 Min mor
Balsari 1999 (98)	NR	NR	NR	NR	2000 rpm	20 min	NR	-20°C	30	230

30–1752 Min–max Median = 294	NR	12–492 Min-max Median = 66	NR	40–530 Min_mav	4–720 Min-max	Median = 180 Median = 250 postmenopausal Median = 104	premenopausar 119.2–282.7 Min–max	232.4 ± 163.6 (Mean + SD)	201.7 mean Median = 167 5	250 (95°perc.) NR	6.01–774.76 Min–max	Median = 261.13 NR	(continued)
145	NR	56	NR	32	63	64	б	42	88	26 NR	20	NR	
-80°C	-80°C	-70°C	-70°C	-20°C	-80°C	-70°C	-80°C	-80°C	-80°C	-80°C	-70°C	-80°C	
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Kraft 1999 (<i>99</i>)	Salgado 1999 (37)	Salven 1999 (43)	Salven 1999 (44)	Wynendaele 1999 (100)	Adams 2000 (40)	Byrne 2000 (29)	Gunsilius 2000 (101)	Lee 2000 (108)	Heer 2001 (74)	Benoy 2002 (45) Lantzsch 2002 (102)	McIlhenny 2002 (50)	Bachelot 2003 (103)	

AnticoagulantTemperatureCentrifugationAuthors (ref.)TypeConc.Image of sampleCentrifugationVerheul 1997 (38)EDTANRRTNRNRNBanks 1998 (39)EDTANRNRRTNRNRNMaloney 1998 (96)EDTANR $0.12-0.2\%$ (w/v)RT $2000g$ 10 min1Webb 1998 (97)EDTANR 7.5% (w/v)RT $2000g$ 10 min2Webb 1999 (43)Sodium citrate $0.12-0.2\%$ (w/v)RT $2000g$ 10 min2Wrendaele 1999 (100)K_1 EDTA 7.5% (w/v)RT $2000g$ 10 min2Wynendaele 1999 (100)K_1 EDTA $0.129M$ NR $3000g$ 10 min2Wynendaele 1999 (100)KaEDTA $0.129M$ NR $3000g$ 10 min2Wore 1000Tisodium citrate 0.31% (w/v)NR $3000g$ 10 min2Wore 1000Tisodium citrate 0.31% (w/v)NR $2000g$ 10 min2Malams 2000 (101)Sodium citrate 0.31% (w/v)NR $NRNRNRNRLee 2000 (108)EDTANRNRNRNRNRNRNRNRMcIlhenny 2002 (50)EDTANRNRNRNRNRNRNRNRWu 2002 (79)EDTANRNRNRNRNRNRNNSodium citrateWu 2002 (79)ED$				·						
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		4	NR	RT	NR	NR	NR	-20°C	30	0-20
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										Median = 30

Table 2 (continued) b. Plasma

276

IQR, Interquartile range.

to 8-10 times (38). So, serum VEGF concentration mainly reflects platelet counts rather than the tumor burden. For this reason, some authors suggested the use of a specific matrix to avoid the potential hazard for systematic errors.

The majority of studies on the role of VEGF as a circulating tumor marker used serum prepared by different procedures: the time from venipuncture to centrifugation was quite variable, ranging from "immediately" to 30 min at room temperature, and from 30 min to 4 h at 4°C. In a study that standardizes the sampling procedure for the determination of VEGF in different blood fractions, Banks et al. (39) demonstrated that in a time range from 0 to 2 h at room temperature, serum VEGF increased from a minimum of 118% to a maximum of 4515%, with respect to the VEGF levels found in the serum obtained 10 min after blood withdrawal. The maximum VEGF released from clotting is reached only after 2 h at room temperature.

Also clinical studies of plasma VEGF levels have been performed using a wide variety of methods for sample collection and handling: as far as the type of anticoagulant is concerned, some authors suggested the use of sodium citrate plasma (40,41). Banks et al. (39) demonstrated that sodium citrate does not prevent activation of platelets, and the authors propose the use of CTAD plasma, it being more effective in preventing the release of VEGF by platelets. VEGF has also been investigated in whole blood (42), while others estimated the theoretical VEGF load of platelets as the ratio of VEGF serum concentration related to the platelets number (39,43–46). The biological significance of white blood cell–derived VEGF has yet to be clarified. To date there is only reported a significant association between whole blood VEGF and leukocyte count. Hence, it remains unclear in cancer patients which measurement of circulating VEGF actually represents the fraction of biologically active VEGF in the circulation.

4.1.2. BIOLOGICAL REASONS FOR VARIABILITY

VEGF levels have been studied also in relation to the menstrual cycle. However, contradictory results have been found with some studies suggesting correlation with hormones, some found higher levels of serum VEGF in the follicular phase (47), some in the luteal phase (48), while others did not find consistent changes (49,50).

In a study Meo et al. (51) concluded that a difference between serial results can be considered as normal if the variation is less than about 60% in the plasma CTAD, 33% in serum and 43% in whole blood. Moreover, the individuality Index (ratio between intrasubject biological variation and intersubject biological variation) has shown that the range of reference used in some studies could be misleading and that the VEGF determination could be more useful in monitoring the variation within a subject, for example, during therapy, considering the found value of critical difference.

Table 3 Ways of Targeting VEGF

Ligand sequestration Attack external membrane receptor Inhibit VEGF-R message Indirect inhibition

4.2. Clinical Studies

How to best target VEGF has been the subject of several investigative approaches, many of which have been applied in the clinic (Table 3). Several studies are being conducted to evaluate the safety profile and activity of the combination of standard chemotherapy with new antiangiogenetic agents. However, definitive results on the effect on angiogenesis blood markers have been reported only in a few of them to date (Table 4).

4.3. Anti-VEGF Therapy

4.3.1. LIGAND SEQUESTRATION

Ligand sequestration has been approached with the use of monoclonal antibodies directed against VEGF, several of which have been generated and clinical trials are underway. A recombinant humanized monoclonal antibody has been developed that recognizes all VEGF isoforms (rhuMab VEGF) without binding to FGF, HGF, PDGF, or nerve growth factor. It has shown to have indirect and potent antitumor activity in experimental models. The safety of bevacizumab was evaluated in a phase I dose-escalating study involving 25 patients who had a variety of tumors, including sarcomas (n = 8), renal cancer (n = 7) and breast cancer (n = 2) (52,53).

Dose-limiting toxicities were not observed at weekly doses of $\leq 10 \text{ mg/kg}$, although some patients experienced asthenia, mild headache, fatigue, nausea, and low-grade fever on the day of the administration. In addition, bleeding at tumor sites developed in 3 (12%) of the 25 patients. No objective response was observed; two patients obtained a minor response. Twelve (52%) of the 23 patents had stabilization of disease during the 70-d study period. As a single agent, rhuMAb VEGF has been shown by Sledge et al. (54) to induce remissions and prolonged stabilization of disease in patients with heavily pretreated metastatic breast cancer. A phase III trial has been conducted in advanced breast cancer patients and the results recently published. This study was performed to evaluate the combination of bevacizumab with capecitabine vs capecitabine alone in patients affected by metastatic breast cancer previously treated with anthracycline and taxane. Patients were randomized to either capecitabine alone or in combination with bevacizumab. Treatment was continued until evidence of progression;

Drug	Mechanism of action	Clinical development
BEVACIZUMAB	Anti-VEGF monoclonal antibody	Phase III: Miller et al. (55)
SU5416	Inhibition of Flk-1 (VEGFR)	Phase I: Stopeck et al. (68)
ZD6474	Inhibition of VEGFR and EGFR	Phase I: Minami et al. (69)
CEP-7055	Inhibition of VEGFR1-2-3	Phase I: Ruggeri et al. (70)
SU6668	Inhibition of tyrosine kinase receptor of VEGF, bFGF, PDGF, c-Kit	Phase I: Hoeckman et al. (71)
PTK 787/ZK 222584	Inhibition of VEGF-R1 (Flt-1) and VEGF-R2 (Flk-1/KDR)	Phase I: Thomas et al. (72)
Angiozyme	Constructs targeting VEGFR Flt-1 mRNA	Phase I–Phase II (105–107)

Table 4 Clinical Studies patients randomized to the combination arm of the study were eligible to receive bevacizumab after progression, either alone or with other therapies. Promising results have been observed in terms of response rate, but no benefit was obtained regarding overall survival (55).

Monoclonal antibodies have also been developed against the external membrane domain of the VEGF-R2 receptor (56,57) and clinical trials have recently been initiated with these agents, but there are no published reports are to date.

4.3.2. Tyrosine Kinase Inhibitors

Several receptor tyrosine kinase inhibitors directed against the internal membrane tyrosine kinase portion of VEGF receptors 1 and/or 2, have been developed (58–65). Many of these compounds target more than one receptor: SU6668 targets the VEGF-R2 as well as the PDGF and FGF-1 receptors; PTK 787/ZK 222584 targets both VEGF-R1 and R2 (63).

SU5416 is a small, lipophilic, highly protein-binding synthetic molecule that inhibits phosphorylation of the selective tyrosine kinase for the VEGF receptor Flk-1 on endothelial cells (66). In addition, it binds to the PFDG receptor, which is also involved in the transduction of angiogenesis signals. Finally, it binds c-Kit, a related tyrosine kinase receptor for stem cell factor and a hematopoietic growth factor that promotes the survival of hematopoietic progenitor cells and in multiple lineages (52,67). SU5416 produces a dose-dependent inhibition of tumor growth in a variety of xenograft models, including malignant melanoma, glioma, fibrosarcoma, and carcinomas of the lung, breast, prostate and skin (52,66). In a human colon cancer xenograft model, SU5416 inhibited tumor metastases, microvessel formation, and cell proliferation (52).

Phase I and II clinical trials have been completed using a twice-weekly dosing regimen with a maximum tolerated dose equal to 145 mg/mq. Stopeck et al. (68) performed a phase I dose-escalating study of SU5416 in 22 patients with advanced malignancies, 3 of them were affected by breast cancer. Of the 19 evaluable patients, 1 obtained a partial regression and 3 obtained disease stabilization for at least 12 wk.

ZD6474 is a novel vascular selective endothelial growth factor receptor tyrosine kinase inhibitor that also has activity against epidermal growth factor receptor tyrosine kinase. A phase I study of ZD6474 has been conducted in 18 Japanese patients with solid tumors refractory to standard therapy. Most patients treated had non-small-cell lung cancer (NSCLC) or colorectal cancer (CRC). A dose range of 100–300 mg/d of ZD6474 was well tolerated and considered to be appropriate in terms of efficacy for use in phase II studies of patients with NSCLC (*69*).

CEP-7055 is a low molecular weight tyrosine kinase inhibitor with activity against VEGF-R1, 2, and 3. In a range of experimental models, CEP-7055 has been shown to induce significant inhibition of both angiogenesis and tumor growth. The clinical activity and tolerability of CEP-7055 has been explored in a phase I study: 19 patients have been recruited at doses of 10, 20, 40, 80, and 120 mg given twice daily. Pharmacodynamic assessments of tumor blood flow by PET, dermal wound angiogenesis, and other blood markers are being performed (70).

SU6668 is a novel compound that competitively inhibits the tyrosine kinase of the receptors of VEGF, bFGF, PDGF, and c-Kit. SU6668 inhibits angiogenesis through several mechanisms, primarily by the induction of apoptosis in both endothelial and tumor cells. Currently, phase I studies are underway to evaluate the potential of SU6668 as anticancer agent for humans (71).

PTK 787/ZK 222584 (PTK/ZK) is an oral potent and selective inhibitor of VEGF-mediated Flt-1 and KDR receptor tyrosine kinases. Phase I studies are underway evaluating the optimum dose and schedule of oral PTK/ZK administered continuously to patients with advanced cancers known to overexpress VEGF. To date, particularly in patients with liver metastases from colorectal cancer treated with PTK/ZK, dynamic contrast-enhanced magnetic resonance imaging has been a useful predictor of the biological response of VEGF-receptor inhibition (72).

At least another five selective anti-VEGF small tyrosine kinase inhibitors are entering clinical evaluation.

4.3.3. HAS VEGF DETERMINATION A PROGNOSTIC-PREDICTIVE VALUE?

Based on the finding that in breast cancer intratumoral VEGF expression and microvessel density significantly correlate with decreased relapse-free survival (73), several studies have been conducted to evaluate the prognostic-predictive value of circulating VEGF in breast cancer (Table 5). The relationship of preoperative serum VEGF with conventional prognostic indicators of breast cancer has been recently studied and serum VEGF was compared with two established tumor markers for breast cancer: CEA and CA 15-3 (74). This prospective study involved 200 patients. Eighty-eight healthy females were also recruited as controls for serum VEGF. This study showed that serum VEGF is significantly high in ductal but not in lobular carcinoma. This study agrees with Dvorak et al. (22), who showed that only lobular carcinoma of the breast and papillary carcinoma of the bladder failed to reveal significant VEGF mRNA expression. Moreover, it has been shown that serum VEGF has a much higher sensitivity (62.1%) in detecting breast cancer than both the tumor markers CA 15-3 (13.6%) and CEA (10.3%), with a specificity of 74%.

A difference in the angiogenic response of different types of breast cancers would allow the selection of patients for whom personalized adjuvant therapy is needed, similarly to the selection of patients for tamoxifen therapy

Study	Results
Serum VEGF levels and tumor markers for breast cancer	VEGF shows higher sensitivity when compared to CEA and CA 15-3 in detecting breast cancer (74).
Plasma VEGF levels and progression of breast cancer	Plasma VEGF is an independent predictor of overall survival of local recurrence (79).
Serum VEGF levels and endostatin	Elevated serum VEGF levels are correlated with short free-relapse survival (81).
Serum VEGF levels and response to chemotherapy	Normalization or decline >50% in VEGF levels is significantly higher in patients who responded to chemotherapy then in nonresponders (83).
Serum VEGF levels and response to thalidomide	VEGF levels changes not significantly correlates to response to thalidomide (87,90).

 Table 5

 Prognostic/Predictive Value of Circulating VEGF in Breast Cancer

on the basis of the ER status. Various factors such as estrogen, protein kinase C, and cAMP can induce the expression of these gene complexes, which in turn can up-regulate VEGF expression (75). It has been demonstrated that the genetic status of a tumor may determine whether or not estrogen can stimulate VEGF expression (76). However, long term follow-up studies are required to determine whether preoperative serum VEGF levels are of prognostic significance, as has been shown for tumor VEGF levels, and whether serum VEGF will be useful in detecting early recurrence in breast cancer patients (74).

Based on the finding that plasma VEGF shows better discrimination than serum VEGF (77) and that no information is so far available on the prognostic relevance of plasma VEGF levels in patients who have had surgery for breast cancer (78), Wu et al. (79) performed a study on plasma VEGF to assess whether this biomarker is associated with progression of breast cancer in African-American and Hispanic women. Wu et al. (79) measured plasma VEGF of 125 women after adjuvant treatments and examined the association of plasma VEGF levels with other tumor characteristics such as steroid hormone receptors, tumor size, regional nodes, and stage. Plasma VEGF levels were found to be significantly higher in breast cancer patients than in normal subjects and in patients with large tumor size, and stage III/IV disease. Multivariate analysis showed that plasma VEGF was an independent predictor of overall survival of local recurrence. These findings suggest that plasma VEGF should be considered as a tumor marker of breast cancer progression.

However, the dispute whether serum VEGF or plasma VEGF should be used as a surrogate biomarker of angiogenesis is still controversial. Some studies suggest that platelet-poor plasma more accurately reflect tumor progression, whereas others found that serum VEGF gives a better prognostic information. On the basis of the hypothesis that activated platelets in tumor vasculature release thrombopoietin and, thereby, stimulate bone marrow generation of platelets, both plasma VEGF and serum VEGF may be important for tumor activity. Plasma VEGF may be increased because of direct tumor release of VEGF, and by tumor-induced intravascular platelet activation and subsequent VEGF release. Elevated serum VEGF may be the consequence of increased platelet numbers in cancer patient, caused by intratumoral platelet activation and subsequent release of thrombopoietin.

Recently, Nishimura et al. (80) have measured plasma VEGF levels in various breast diseases. Fifteen patients had benign breast disease, 187 patients primary breast cancer: 32 patients of whom without postoperative recurrence, and 56 patients with recurrence. Plasma VEGF levels were measured by enzyme-linked immunosorbent assay (ELISA). Plasma VEGF levels were higher in malignant than in benign breast disease, and were also higher in patients with recurrence or distant metastasis.

More recently, Zhao et al. (81) also investigated the correlation between serum VEGF and endostatin levels in patients with breast cancer. Serum VEGF and endogenous endostatin levels were detected before surgery and 3 wk after surgery. Preoperatively, the levels of the two biomarkers were significantly elevated and correlated with each other. Postoperatively, VEGF levels decreased significantly while those of endostatin remained high. Patients with both normalized VEGF and elevated endostatin following surgery had a lower risk of relapse. Univariate and multivariate analysis showed a correlation between elevated VEGF level and short free-relapse survival.

A similar study has been conducted to investigate whether increased VEGF serum levels correlate with poor outcome in advanced colorectal cancer patients. Catalano et al. (82) investigated in a retrospective study the pretreatment serum levels of VEGF and their correlation with outcome in 140 consecutive colorectal cancer patients and 50 healthy subjects. Fifty-seven patients were staged as locally advanced disease (stage III), while 83 had metastatic disease (stage IV). The median pretreatment VEGF serum levels were significantly higher in colorectal cancer patients as compared to control subjects. Median VEGF serum levels were shown to be significantly higher in stage IV patients than in those with stage III (834 pg/mL vs 435 pg/mL, when basal VEGF serum levels in metastatic patients were

analyzed according to response to chemotherapy, responders patients were shown to have significantly lower VEGF serum levels than non-responders: 398 pg/mL vs 734 pg/mL. In univariate analysis basal VEGF serum levels correlated with both overall survival and time to treatment failure. A multivariate analysis included pretreatment serum VEGF levels, ECOG performance status and tumor stage as the independent prognostic factors for overall survival and time to failure. These results suggest that the evaluation of pretreatment serum VEGF levels may be useful for predicting outcome in patients with locally advanced or metastatic colorectal cancer.

The VEGF signaling pathway provides many different levels or targets that could be used to modify or disrupt the transmission of the signal, both upstream or downstream of VEGFR binding. Since VEGF represents the main angiogenic factor, the control of VEGF secretion could represent the most important target for inhibition of angiogenesis-related tumor growth.

In experimental studies, certain chemotherapeutic agents such as paclitaxel inhibit VEGF-induced angiogenesis, while at present there are no data on the possible influence of chemotherapy on VEGF secretion in cancer patients. A preliminary study was conducted by Lissoni et al. (83). The study included 14 patients with metastatic breast cancer treated with paclitaxel. Serum levels of VEGF were measured by ELISA in blood samples collected before and after therapy and at 21-d intervals. The clinical response consisted of partial response (PR) in three and stable disease (SD) in six patients. The percentage of normalization or decline greater than 50% in VEGF levels was significantly higher in patients with PR or SD than in those with progressive disease (PD) (5/9 vs 0/5). This preliminary study suggests that the efficacy of paclitaxel, at least in terms of disease stabilization, may be associated with decreased VEGF blood levels of course, the small size of the study prevents any definitive conclusion.

VEGF is commonly over-expressed in pancreatic cancer and its expression appears to be an important predictor of survival. At the 39th ASCO meeting a study was presented of the combination of bevacizumab plus gemcitabine in patients with advanced pancreatic cancer. Twenty-one patients were enrolled to receive gemcitabine at a dose of 1000 mg/m² given intravenously on d 1, 8, and 15, every 28 d and bevacizumab at a dose of 10 mg/kg given intravenously d 1, 15. Pretreatment serum VEGF levels were collected. The study is still ongoing and 16 patients are currently evaluable for response in this trial. There have been 6 confirmed partial responses (38%), 7 patients (44%) had stable disease. Median survival has not been reached. Median time to progression is 5.5 mo. Pretreatment VEGF levels range from 0 to 586 pg/mL and do not correlate with clinical end points. Estimated 1-yr survival is 54% (84).

Bevacizumab has also been tested in patients with advanced melanoma in a study by Carson et al. (85). Patients with metastatic melanoma who had not received prior cytokine therapy were eligible. They were randomized to receive bevacizumab (15 mg/kg intravenously every 2 wk) \pm low dose α -interferon (1 million units/m² daily). Patients were restaged after 12 wk and received 12 more wk of therapy if they had a clinical response or stable disease. There were two responses in the combination arm. Four patients (three bevacizumab, one bevacizumab+interferon) had prolonged stabilization of disease. Baseline levels of VEGF were measured by ELISA and ranged from 471 to 2686 pg/mL except for one patient who had levels of >4000 pg/mL. The above patient had the longest period of stabilization of disease. Otherwise, VEGF levels did not correlate with response. By contrast, FGF levels ranged from 0 to 180 pg/mL at baseline and dropped significantly during cycle 1 in the two response patients and two of four SD patients. The accrual is ongoing.

Interesting results have been observed with the use of thalidomide, a drug developed as a sedative, that has shown antitumor activity by various mechanisms of action: inhibition of angiogenesis, cytokine-mediated pathways, modulation of adhesion molecules, inhibition of cyclooxygenase-2, and stimulation of immunoresponse (86). Baidas et al. (87) performed a randomized phase II study of thalidomide in patients affected by metastatic breast cancer: 28 patients were randomized to receive either 200 mg/d (arm A) or 800 mg/d (arm B) to be escalated to 1200 mg. No partial or complete responses were observed. Two patients in arm A had stable disease. Changes in serum bFGF, VEGF, and TNF- α levels from baseline to time of removal from study in 26 patients were determined. The results of these analysis indicate mean percentage changes from baseline of -37% for bFGF, +60% for VEGF, and +79% for TNF- α . Of these, only the increase in TNF- α levels was statistically significant.

The vascular pattern of renal cell carcinoma (RCC) suggests that angiogenesis inhibition may be a new biological treatment for patients with this disease (88,89). Several studies have been conducted testing thalidomide in metastatic RCC (MRCC), Minor et al. (90) evaluated the toxicity and activity of thalidomide and measured the changes of VEGF 165 plasma levels after therapy. In this phase II study, 29 patients were enrolled and the drug was given using a intrapatient dose escalation schedule, starting at a daily dose of 400 mg and escalated as tolerated up to 1200 mg. Of the 24 patients evaluable for response only 1 partial response, 1 minor response, and 2 stable disease for over 6 mo were observed. VEGF 165 levels were not modified by therapy.

Endostatin, a 20-kDa fragment of collagen XVIII, is an endogenous angiogenesis inhibitor that has been shown to inhibit, in a potent and dosedependent manner, the growth of a wide variety of human and murine primary metastatic tumors in mice (91,92). Herbst et al. (93) have evaluated recombinant human endostatin (rh-Endo) in a phase I study designed to assess safety, pharmacokinetics, and serum markers of angiogenesis in patients with refractory solid tumors. The serum levels of VCAM-1 were assayed before the study and every 28 d, using the ELISA method. Twenty-five patients were treated and a considerable variation in the baseline levels of all four factors was observed. Furthermore, after of therapy with rh-endostatin, there were no consistent changes in the levels of any of these proteins. The absence of a measurable effect on VEGF, bFGF, VCAM, and E-selectin also included the two patients who experienced minor anticancer effects (93).

5. CONCLUSIONS

Determination of circulating VEGF still remains an experimental procedure with no evident application for routine clinical decisions. Data from retrospective studies, however, suggest that VEGF levels may predict clinical outcome of breast cancer. Prospective studies are needed to properly evaluate the method of choice and the clinical significance of preoperative determination of tissue or circulating VEGF for prognostic purposes. Only a minority of phase I studies with selective inhibitors of angiogenesis evaluated the biological activity as antiangiogenic compounds by determination of surrogate biomarkers. Therefore, both the selection of the patients and the prediction of response to anti VEGF therapy still remain unanswered questions. From a biological point of view the pathways regulating tumor hypoxia and VEGF expression seem to be highly correlated, so codetermination of both may allow for more accurate information on the angiogenica status of each single tumor. Clinical prospective studies should be planned to test the value of surrogate markers of angiogenesis as an integral part of the studydesign for the testing of a new antiangiogenic compounds.

Properly designed projects of translational research aimed to correlate information from laboratory assays with activity of new therapeutic strategies based on the use of biological response modifiers/molecular-targeting agents should also foresee the use of adequate methods of validation of laboratory methods, with procedures of standardization and quality controls. Recent clinical data on the activity of bevacizumab associated with chemotherapy in colorectal and renal cancers in unselected series of patients are promising, also regarding the expression of the molecular target. The discovery of predictive biomarkers for response are presumed to be the key to improve the selection of the cases to be treated with anti-VEGF compounds in future studies.

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15 Detection of Early Tumor Dissemination in Patients With Breast Cancer

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CONTENTS

INTRODUCTION DETECTION OF OCCULT METASTASES IN THE PERIPHERAL BLOOD DETECTION OF OCCULT METASTASES IN THE BONE MARROW (OMBM) DETECTION OF OCCULT METASTASES IN THE LYMPH NODES OCCULT METASTASES IN THE BONE MARROW AND LYMPH NODES FUTURE DIRECTIONS ONGOING PROSPECTIVE STUDIES PATIENT-SPECIFIC TREATMENT REFERENCES

SUMMARY

Women who are newly diagnosed with breast cancer have operable disease, and therefore are potentially curable. Nevertheless, a subpopulation of these patients, who are thought to be disease free after initial treatment, go on to develop recurrent disease. These recurrences are due to early spread of tumor cells, either systemically

Cancer Drug Discovery and Development: Biomarkers in Breast Cancer: Molecular Diagnostics for Predicting and Monitoring Therapeutic Effect Edited by: G. Gasparini and D. F. Hayes © Humana Press Inc., Totowa, NJ (peripheral blood and/or bone marrow) or regionally (lymph nodes) that are not detected by methods routinely employed. To address this problem, more sensitive methodologies of detecting early disseminated tumor cells have been developed over the past decade and a half. This chapter looks at the more important methods and discusses the clinical relevance of these methods.

Key Words: Bone marrow; circulating tumor cells; early tumor dissemination; flow cytometry; immunohistochemistry; immunomagnetic separation; lymph nodes; micrometastasis; occult metastasis; peripheral blood; prognosis; reverse transcriptase-polymerase chain reaction; sentinel lymph nodes.

1. INTRODUCTION

As with most types of cancer, early detection and definitive treatment are the keys to disease-free survival in patients with breast cancer. The majority of patients with newly diagnosed breast cancer have operable disease, and these patients are considered potentially curable. The traditional prognostic indicators for patients with operable disease are the size of the primary tumor and the spread of disease, either regionally to the lymph nodes or systemically to the bone marrow. However, up to 35% of patients, including up to 7–19% of patients with no evidence of metastasis at the time of diagnosis develop recurrent disease after primary therapy (1,2). This rate of disease recurrence in patients with no evidence of regional or systemic metastasis at the time of initial treatment supports the notion that traditional methods are not sensitive enough to detect the earliest metastatic spread of disease, as clearly, these patients had occult systemic spread of disease that was undetectabled. As a result of this, several groups have recommended adjuvant treatment for patients with lymph node negative disease (1-6). While this is controversial (because the majority of node negative patients will be clinically cured without adjuvant therapy), it is in this group of patients who have early disseminated tumor cells either regionally or systemically, that adjuvant therapy should be most successful.

A substantial amount of work has been done over the past decade and a half to develop more sensitive, reproducible and clinically relevant methods of detecting the earliest evidence of tumor cell dissemination (7). Several terms denoting these early-disseminated tumors cells are currently used, most commonly, micrometastases (MMs), occult metastasis (OM), occult micrometastasis (OMM), minimal residual disease (MRD), circulating epithelial cells (CECs), and early-disseminated tumor cells (EDTCs). We will use the term occult metastasis (OM).

This discussion focuses on the detection and significance of occult metastatic tumor cells in the peripheral blood, bone marrow (OMBM), and lymph nodes (OMLM) of patients with breast cancer as well as on the biologic characterization of these cells and potential for the development of specific molecular targets.

1.1. Strategies Employed for the Detection of Occult Metastases

Several techniques have been used to identify occult metastases in the various compartments. The most important of these include immunohis-tochemistry (IHC), flow cytometry, and molecular methods, usually the reverse-transcriptase polymerase chain reaction (RT-PCR) technique. Each method has its advantages and disadvantages. We will discuss each of the major techniques commonly in use today.

1.1.2. FLOW CYTOMETRY

A flow cytometric assay was developed to detect rare cancer cells in the blood and bone marrow (8). The method has been reported to be extremely sensitive, with an ability to detect as little as one positive cell in ten million blood cells in a model system, although this level of sensitivity has not been universally obtained (9). Leers and associates (10) assessed the utility of multiparameter flow cytometry (MP-FCM) using an anticytokeratin antibody, to determine whether or not it is an effective method for the detection of occult metastases in the lymph nodes in patients with breast cancer. Of the 38 cases they examined in which the lymph nodes were positive by hematoxylin and eosin (H&E) and/or IHC they were able to detect more than 1% cytokeratin-positive cells in 37 cases. They concluded that MP-FCM was a highly specific and sensitive nonmorphological technique and provided several advantages including: (1) almost the entire lymph node is assessed, (2) the technique is able to provide information concerning ploidy, and (3) it may be possible to introduce sorting of metastatic tumor cells for further molecular studies. One major disadvantage of most flow cytometric systems has traditionally been the inability to morphologically characterize the cells constituting the "positive" events. However, by employing sophisticated cell-sorting technologies, in which the extrinsic cell population can be captured for subsequent morphologic evaluation, the specificity of tumor cell detection might be improved.

1.1.3. IHC

IHC was one if the earliest techniques employed for the detection of occult metastases and remains the most important method to date. Pioneering studies at the Ludwig Institute and Royal Marsden Hospital in London, England (11), helped to establish specific IHC procedures to identify occult metastatic cancer cells in the peripheral blood and bone marrow of patients with cancer. The initial focus was on the study of breast cancer (11–15),

although many other tumor types have subsequently been studied including colon (16-19), prostate (20-23), lung (24-32), esophagus (33-37), and melanoma (38-40). IHC methods are based on the ability of monoclonal antibodies to distinguish between cells of different histogenesis (i.e., epithelial cancer cells vs the hematopoietic cells of the peripheral blood, bone marrow, and lymph nodes). The results indicate that it is possible to identify occult metastatic cancer cells in these compartments prior to their detection by routine histologic analyses, and that the presence of these cells may be an important risk factor for disease recurrence.

The most widely used antibodies to detect occult metastatic tumor cells are monoclonal antibodies to epithelial-specific antigens. These antibodies do not react with hematopoietic cells normally present in the peripheral blood, bone marrow, and lymph nodes. None of the antibodies used in any study is specific for cancer; all react with normal and malignant epithelial cells. They are useful because they can identify an extrinsic population of epithelial cells in the blood and bone marrow, where there are normally no epithelial elements. The reported sensitivity of the IHC method ranges from the detection of one epithelial cell in 10,000 (41) to that of two to five epithelial cells in a million hematopoietic cells (14,41,42).

Because current methods used to detect occult metastases generally involve epithelial markers that are not specific for malignant cells, concerns have been raised over interpreting artifactually present epithelial cells as tumor cells (43). Certainly, keratin-positive epithelial cells have been found on the surface of "clean" slides and floating on water baths and in staining solutions. Perhaps a greater danger is the possibility that tumor cells may be mechanically shed from the primary tumor to the lymph nodes as a result of manipulation of the tumor, for example as a result of fine needle aspiration or core biopsy. The danger exists that these cells could be overinterpreted as occult metastatic cells by an inexperienced observer. However, normally exfoliated epithelial cells lack the morphologic features of tumor cells and can be differentiated from metastatic malignant cells on the basis of size, nuclear hyperchromasia, nuclear to cytoplasmic ratio, and location on the slide by an experienced pathologist. In addition, the number of positive events and their pattern of distribution in the lymph node can give valuable clues as to their origin.

1.1.4. MOLECULAR METHODS

While IHC methods are effective in identifying occult metastases in the peripheral blood, bone marrow and lymph nodes of patients with operable breast cancer, these methods are laborious and require considerable technical expertise to perform and interpret. Efforts are under way to develop molecular techniques for the detection of occult metastases. The method usually employed is RT-PCR, which differentiates gene expression between

epithelial and lymphoid cells to identify epithelial cancer cells. RT-PCR entails the isolation and reverse-transcription of epithelial-specific messenger RNA to complementary-DNA (cDNA), and thereafter, involves PCRbased amplification of the cDNA template between specific primers. This results in a several thousand-fold amplification of the signal, and makes the method theoretically extremely sensitive. Keratin 18 or 19 (CK-18, CK-19) mRNA that is expressed in epithelial cells has been the most frequent target for amplification to detect occult metastases. Studies have (44) used the CK-19 transcript to identify cancer cells in the bone marrow and peripheral blood of breast cancer patients. However, it is known that a pseudogene for CK-19 exists that shares a very high homology with the mature mRNA from the CK-19 gene. Contamination of DNA in the RNA sample used for reverse transcription can result in amplification of the pseudogene, giving rise to specific "background" bands in the negative controls (known negative bone marrow and blood) (23). Furthermore, the nonepithelial cells may have a low level of expression of epithelial transcripts, leading to background bands. To evaluate the specificity and sensitivity of RT-PCR in detecting occult metastases, transcripts were amplified from epithelial and breast cellspecific genes including carcinoembryonic antigen (CEA), cytokeratin 19 (CK-19), cytokeratin-20 (CK-20), mucin-1 (MUC-1), and gastrointestinal tumor-associated antigen 733.2 (GA733.2), from the blood of healthy donors and lymph nodes from patients without cancer by RT-PCR. CK-20 was the only marker not detected in the lymph nodes or blood from patients without cancer (45). Similarly, the specificity of RT-PCR assays was studied (46) using primers specific for various tumor-associated and organ-specific mRNA species including prostate-specific antigen (PSA), epithelial glycoprotein-40 (EGP-40) desmoplakin I (DPI I) CEA, erb-B2, erb-B3 prostatespecific membrane antigen (PSM), and CK-18. The bone marrow from 53 subjects with no epithelial malignancy as well as bone marrow samples from 53 patients with prostate cancer and 10 patients with breast cancer were examined. It was found that seven of the eight markers tested could be detected in a considerable number of bone marrow samples from control patients. In this study only PSA mRNA was not detected in any of the 53 control bone marrow samples (46).

The known relative nonspecificity of epithelial cell markers has prompted the search for even more specific breast cancer markers. One such candidate currently under investigation is mammaglobin (hMAM). Mammaglobin, a glycoprotein expressed in the mammary glands of women, as well as in breast cancer cell lines, has been investigated as a potential marker to detect early metastatic breast cancer in the peripheral blood (47–49) of patients with breast cancer. The specificity and sensitivity of hMAM will be discussed in detail later. In summary, blood samples from patients with breast cancer and other malignancies tested for hMAM by RT-PCR showed the marker to be relatively sensitive but lacked specificity when analyzed alone (47).

Metastatic melanoma has been successfully detected by RT-PCR using mRNA markers for tyrosinase, p 97 and MelanA/MART1.

In summary, the drawbacks of molecular methods include the chance of low level of epithelial gene expression from lymphoid cells that could result in high background, as well as the inability to employ morphologic criteria to confirm the presence of metastatic cells. However, RT-PCR has been successfully used for the detection of occult metastases to the bone marrow in melanoma and carcinoma of the prostate and colon.

Techniques to detect occult metastases are neither easy to perform nor simple to evaluate. We therefore, recommend that these tests be preformed by laboratories that are experienced in doing large numbers of them and that they be evaluated by specialists who are thoroughly trained in their interpretation.

2. DETECTION OF OCCULT METASTASES IN THE PERIPHERAL BLOOD

Clearly a technique that could reliably detect the presence of early tumor in the peripheral blood would be a great practical advantage over one in which a sample of bone marrow was required. There have been a number of studies designed to explore this possibility. Unfortunately, the yield of occult metastatic cells from peripheral blood is extremely low. Redding et al. (11) found that 28.2% patients with breast cancer showed extrinsic cancer cells in their bone marrow, but only 2.7% of these patients had detectable cells in their peripheral blood. It is not clear why tumor cells are detected less commonly in peripheral blood than in bone marrow. For this reason most of the studies looking for the presence of early metastases in the blood have focused on using molecular methods. Nevertheless important work continues to examine occult metastases in the peripheral blood by a variety of methods and markers especially in the area of breast cancer.

2.1. IHC vs RT-PCR

To compare the sensitivity of OM in blood by molecular and IHC methods, both peripheral blood and bone marrow samples from women with primary breast cancer using RT-PCR and IHC have been analyzed (50). The study demonstrated that the peripheral blood is a less sensitive compartment than the bone marrow (9% vs 43%) (51). The reason for this difference in sensitivities is not known. It has been suggested that this lack of sensitivity in detecting tumor cells in the blood may be, at least in part, because tumor cells are likely to only be shed intermittently into the circulation (51). Therefore, the lack of sensitivity in blood is a real reflection of the biology of tumor dissemination and not a failure of the detection methods used. The rates of sensitivity of detecting occult metastasis in the blood using RT-PCR vs IHC methodologies was also compared. There was only a small gain in sensitivity using RT-PCR over IHC (51). In summary, it appears that while the methods may not detect an identical population of positive patients and that molecular detection methods may be slightly more sensitive, the number of circulation tumor cells in the blood is less than in the bone marrow.

2.2. RT-PCR

The studies referenced above all examined the peripheral blood for cytokeratin transcripts and/or protein. Molecular (RT-PCR) detection of other markers have been tested in an attempt to improve the sensitivity and specificity of the detection of OM in the blood. Among the most well studied and promising of the new markers are mammaglobin (hMAM) described in Section 1.1.4., and epidermal growth factor receptor (EGFR) (47,49,52). One study found 25% positivity for hMAM in the blood from patients with breast cancer, and in 5% of patients tested with malignancies other than breast cancer, indication that the marker is probably sensitive, but not entirely specific for breast cancer. As many as 43% of patients with metastatic breast cancer who were positive for hMAM by RT-PCR in the peripheral blood have been detected (52). Variations in rates of positivity have been found using the different markers. In a comparison of hMAM, EGFR, and CK-19 in the blood by RT-PCR from patients with breast cancer the rates of positivity were found to be 8%, 10%, and 48%, respectively (47). It was also determined that there is a lack of specificity in some of the markers. None of the blood samples from patients without cancer was positive for hMAM, while CK-19 mRNA was found in the blood of 39% healthy volunteers and transcripts for EGFR and CK-19 were detectable in 25% and 10%, respectively, of the patients with hematological malignancies (47). The lack of sensitivity and specificity found in the markers may indicate that a panel of markers (i.e., CK-19, hMAM, EGFR) may yield superior results than any single marker. Further studies with clinical follow-up is required to test this hypothesis.

2.3. Immunomagnetic Separation

Another method currently under investigation for detecting OM in the blood is the use of immunomagnetic separation techniques. This technology uses antibodies to separate and concentrate the epithelial cells from the hematopoietic cells to improve detection sensitivity. Several enrichment methods are currently available. By the use of various antibody-coupled magnetic particles, tumor cells have been enriched by several orders of magnitude in model tests (42,53). Enrichment can be achieved by positive

or negative selection. Tumor cells can be selected with beads coated with antibodies against tumor-associated antigens (positive selection), or normal blood cells in the preparation can be depleted by use of beads coated with antibodies against hematopoietic cell antigens (negative selection) (54-58). The peripheral blood samples from 21 patients with primary operable breast cancer, 29 patients with metastatic breast cancer, and 21 healthy women have been analyzed by IHC following immunomagnetic separation (59). Tumor cells were not detected in either the healthy women with operable breast cancer. Conversely, 28% women with metastatic breast did have tumor cells in their peripheral blood (59).

2.4. Prognostic Significance of OM in the Blood

The goal of detecting OM in the blood is to us the results to predict patient outcome and develop more patient-specific treatment regimens. Stathopoulou et al. (60) looked at the peripheral blood for the presence and prognostic significance of CK-19-positive cells in patients with breast cancer using the nested RT-PCR assay for CK-19 mRNA. Detection rates for CK-19 mRNA-positive cells in the blood of patients with early or metastatic breast cancer were 74% and 52% respectively. For stage I and II breast cancer, detection of CK-19-positive cells in the peripheral blood before adjuvant therapy was associated with reduced disease-free survival (p =0.0007) and overall survival (p = 0.01). Finally, the hypothesis that the number, not just the presence of circulating tumor cells, was tested using fluorescently labeled monoclonal antibodies cytokeratins (61) from peripheral blood samples from patients with breast cancer. It was found that the levels of circulating tumor cells were significant predictors of progressionfree overall survival (61). These data support the hypothesis that OM in the peripheral blood may be prognostically important

In summary, because of the advantage of ease of accessibility of obtaining the peripheral blood over bone marrow, efforts are underway to optimize currently available detection techniques and to develop new methods of assessing the blood for the presence of early-disseminated tumor. To date, while progress has been made, the peripheral blood continues to be a less sensitive compartment for the assessment of OM than the bone marrow (or lymph nodes). Several markers have been used to detect OM in the peripheral blood of patients with breast cancer including CK-19, hMAM, and EGF-R While none of the markers is entirely specific for the detection of metastatic breast cancer, and the sensitivity of the peripheral blood continues to be less than that of the bone marrow, there is growing evidence that the detection of OM cells in the peripheral blood has a negative impact on prognosis. New techniques such as immunomagnetic cell separation may have an impact in future, but current available data are too preliminary and no clinical follow-up data is available. Further studies are needed with longterm clinical follow-up to determine the clinical utility of OM detection in the peripheral blood.

3. DETECTION OF OCCULT METASTASES IN THE BONE MARROW (OMBM)

The bone marrow vasculature consists of a unique sinusoidal system that may simply act as a filter that traps or concentrates malignant cells. The marrow environment may provide a more favorable support system for tumor cell proliferation than does the blood. One may also speculate that cancer cells that are released into the systemic circulation represent a small subpopulation of cells with altered expression of cell adhesion molecules. Whatever the reason, bone marrow appears to offer the maximum opportunity to detect cancer cells that have been released into the blood. Therefore, the majority of studies to detect occult metastatic cells in the systemic circulation have investigated bone marrow as a site of spread.

Bone marrow in general is the single most common site of breast cancer metastasis, and up to 80% of patients with recurrent tumors will develop metastatic lesions in the bone marrow at some point during evolution of their disease (62); it is also the most frequent initial site of clinically detectable breast cancer metastasis (63). Tumor cells are estimated to be present in the bone marrow of 10 to 45 percent of patients with primary operable breast cancer (11,14,15,64–66), and 20% to 70% of patients with metastatic breast cancer (67). As with most cancers, the most widely used method to detect occult metastatic cells is IHC.

Because IHC is the most widely used (and currently, the most reliable) method for the detection of occult metastases in the bone marrow (OMBM) in breast cancer patients, we have summarized the results from several groups performing IHC assays using monoclonal antibodies.

The percentage of patients with early stage breast cancer in whom extrinsic cells were detected in the bone marrow ranges from 10% to 45%. The possible reasons for some of the variations observed include: (1) use of single antibodies to detect extrinsic cells in some of the studies; (2) differences in patient populations, although all the results were from patients with early stage disease; (3) differences in the antibody reactivity with breast cancer cells; and (4) the presence of antigenic heterogeneity. However, what is evident and striking is that occult metastases in the bone marrow were detected in all of the studies. Furthermore, it is becoming apparent that the rate of detection of occult metastases in the bone marrow in patients with operable breast cancer is decreasing when appropriate antibodies are used. In recent years we have noticed a marked decline in the number of occult metastases–positive cases. We believe that this may, at least in part, be due to the stage migration that has occurred. With the success of early breast cancer detection methods have enabled women to be diagnosed sooner. The latest figures show more than 90% of breast cancer deaths are now diagnosed at a local or regional stage when 5-yr survival rates are 97% and 79%, respectively (68,69). These patients will thus be expected to have a low rate of occult metastases.

Because methods of detecting occult metastases do not specifically target tumor cells, there has been some controversy over whether or not malignant cells are being detected. To help to answer this question, a study was performed by Fehm et al. (70) in which they looked at the blood and primary tumor from patients with cancer (breast, kidney, prostate, and colon) and who had known circulating epithelial cells identified by anticytokeratin antibody. These were analyzed by enumerator DNA probes for chromosomes 1, 3, 4, 7, 8, 11, or 17 by fluorescence *in situ* hybridization and compared to the primary tumor. The pattern of aneusomy matched a clone in the primary tumor in 10 of 13 patients, indicating that the vast majority of these circulating tumor cells were anuesomic and derived from the primary tumor.

3.1. Detection of Occult Metastases in the Bone Marrow in Patients with Early Stage Breast Cancer: Clinical Significance

OMBM have been correlated with known predictors of prognosis in several studies. In our studies (14), the lymph node–negative group had fewer extrinsic cells than the lymph node–positive group, suggesting a trend toward a greater metastatic tumor burden in patients with lymph node metastases.

The presence of OMBM has been correlated with pathologic Tumor– Node–Metastasis (TNM) stage in breast cancer. Several other investigators have obtained similar results. In the original study from the Ludwig Institute (11), the presence of OMBM was correlated with the tumor stage (p = 0.05), and vascular invasion (p < 0.01), both of which are known predictors of poor prognosis.

While the presence of OMBM appears to be correlated with known features of disease progression, the ultimate utility of this test will be determined by whether it can predict breast cancer recurrence. Several studies now show that, in fact, the presence of occult metastases in the bone marrow identifies a population of patients at high risk for recurrence (Table 1) (14,71–76). Our own studies (14) have revealed that the presence of occult metastases in the bone marrow significantly predicts recurrence; the 2-yr recurrence rate for patients with no evidence of OMBM was 3% compared with 33% for patients with detectable OMBM. Diel and associates (71) studied 727 patients with primary operable breast cancer. They were able to detect tumor cells in the bone marrow of 55% of lymph node-positive

Detection of Occult Bone Marrow Metastases in Breast Cancer							
	No. of patients	Clinical follow-up ^a	OMBM [_] (number)	OMBM+ (number)	p-value		
Dearnaley 1991 (77)	39	9.5	31% (8/26)	85% (11/13)	< 0.05		
Mansi 1991 (78)	350	2.3	25% (64/261)	48% (43/89)	< 0.05		
Cote 1991 (14)	49	2^b	16% (5/31)	54% (7/13)	< 0.04		
Diel 1992 (80)	211	2	3% (4/130)	27% (22/81)	0.0001		
Diel 1996 (79)	727	3	8% (34/412)	35% (109/315)	< 0.001		
Braun 2000 (73)	552	4	8% (28/353)	40% (79/199)			
Gebauer 2001 (75)	393	6	20% (46/227)	35% (59/166)	< 0.001		
Gerber 2001 (74)	554	4.5	9%	24%	0.0001		
Wiedswang 2003 (76)	817	4	9% (60/709 ^c)	26% (28/108 ^c)	< 0.001		

Table 1 Detection of Occult Bone Marrow Metastases in Breast Cancer

^aYears.

^bEstimated 2 yr of recurrence rate.

^cOverall survival.

patients and in 31% of lymph node negative patients. They found that OMBM was an independent prognostic indicator for both distant diseasefree survival and overall survival that was superior to axillary lymph node status, tumor stage, and tumor grade. In a more recent study, Mansi and associates (72) looked at bone marrow aspirates from 350 women with primary breast cancer. While they found that patients with OMBM did have shorter relapse-free and overall survival times, it was not found to be an independent prognostic indicator when tumor size, lymph node status and vascular invasion were taken into account (72). Braun and associates (73), analyzed the bone marrow samples from 552 patients with stage I, II, or III breast cancer by IHC. They found cytokeratin positive cells in 36% (199/552) of the patients studied. Forty-nine of the 199 (25%) patients with OMBM died of cancer-related causes, while only 49 of 353 (14%) patients without OMBM died (p < 0.001). They concluded that OMBM was an independent prognostic indicator of the risk of death from cancer (73). Gebauer and associates (75) examined the bone marrow from 393 patients with primary breast cancer. They were able to detect the presence of epithelial cells in 42% of these bone marrow samples. After a median follow-up of 75 months they found that 27% of all patients in the study developed distant metastases. Multivariate analysis showed that, in this study, the presence of bone marrow occult metastasis was an independent prognostic parameter of tumor recurrence or cancer-related death. Gerber and associates (74) looked at 484 patients with pT1-2N0M0 breast cancer. Occult tumor cells were detected in 37.2% of patients, of these 26% had positive bone marrows only, 6.4% had positive lymph nodes only and 4.8% had tumor cells found in both compartments. Patients who had occult metastases in and any compartment showed a decreased disease-free survival and the overall survival rates were found to be comparable top those patients who had one positive lymph node. It appears that occult metastatic tumor cells detected by simultaneous IHC analysis of axillary lymph nodes and bone marrow demonstrate independent metastatic pathways. Finally, Wiedswang et al. (76) examined the bone marrow aspirates from 817 patients with breast cancer. OMBM were detected by IHC using anticytokeratin antibodies. OMBM were detected in 13.2% of all patients and were associated with poor prognosis in node-positive patients and in node-negative patients not receiving adjuvant therapy.

These large studies provide strong evidence that the detection of occult metastases by IHC methods is prognostically important. The presence of OMBM predicts a higher risk for recurrence in bone as well as in other sites. Of particular importance in all of these findings is the fact that the presence of OMBM identifies patients with node-negative disease who are at a higher risk for recurrence; this subset of patients can therefore be the target of more aggressive adjuvant therapy. The studies by both Diel (71) and Braun (73) have shown that bone marrow metastasis is a more powerful predictor of recurrence than histologic node status.

In fact, several studies have now shown that the bone marrow status can be combined with other prognostic factors such as axillary lymph node status. This allows groups of patients to be stratified (14,26,71,77,78) as follows: (1) those with very low recurrence rates (lymph node negative, bone marrow negative); (2) those with moderate rates of recurrence (lymph node negative, bone marrow positive, and lymph node positive, bone marrow negative); and (3) those with high recurrence rates (lymph node positive, bone marrow positive) (14,78-80).

3.1.1. DETECTION OF BONE MARROW OCCULT METASTASES: EFFECT OF TUMOR BURDEN

The number of carcinoma cells detected in the bone marrow (the bone marrow tumor burden) may be significantly associated with disease recurrence (14). In our study, the bone marrow aspirates were processed so that the concentration of bone marrow elements was equal for each patient. Consequently, the number of extrinsic cells counted for each case could be compared among patients; the number of extrinsic cells identified in the bone marrow was considered to be reflective of the peripheral tumor burden in each patient. Among patients with OMBM, those who did not recur had on an average, fewer extrinsic cells in their marrow than those who recurred (15 vs 43 cells respectively). In addition, the estimated 2-yr recurrence rate of patients with 10 or more cells (46%) was significantly higher than that of patients with fewer than 10 cells. Further, the number of extrinsic cells detected in the bone marrow was an independent predictor of prognosis.

4. DETECTION OF OCCULT METASTASES IN THE LYMPH NODES

Despite the explosion on our knowledge of the biology of cancer, the single most important prognostic factor for most solid tumors is the presence of histologically detectable regional lymph node metastases: patients with tumors that have not metastasized to the regional lymph nodes tend to do far better than patients with lymph node metastases. A significant proportion of node-negative patients will, however, develop distant metastasis. As we have seen, systemic dissemination takes place by routes other than lymphatic spread; the presence of bone marrow occult metastases in node-negative patients with. Nevertheless, a proportion of node-negative patients this. Nevertheless, a proportion of node-negative patients without the evidence of bone marrow metastases will experience recurrence. It has now become clear that another possible site for occult tumor spread in histologically node-negative patients is the regional lymph nodes.

Routine histopathological examination of lymph nodes is in reality only a lymph node sampling; in fact, it has been calculated that a pathologist has only a 1% chance of identifying a metastatic focus of cancer with a diameter of three cells in cross-section occupying a lymph node (81). It has also been clearly shown that reexamination of lymph node sections initially considered negative for tumor after routine histopathologic screening frequently shows metastatic deposits, demonstrating that even when tumor cells are present in the section, they can be missed (82). It is evident that routine processing and histologic examination of regional lymph nodes is inadequate to detect the presence of tumor in all cases.

Most studies have involved the detection of regional (axillary) lymph nodes occult metastases in patients with breast cancer (82–89). These studies can be classified into two major categories: (1) detection of metastasis after more intensive histological examination of the lymph node, including analysis of multiple serial sections and (2) studies which use IHC to take advantage of the differential expression of antigens between normal lymph node constituents and epithelial carcinoma cells in order to detect occult tumor in lymph nodes. In fact, this is the same principle as that used for detecting occult tumor in bone marrow. Attempts have also been made to use molecular techniques to detect lymph node occult metastatic cells (89–102).

4.1. Detection of Occult Metastases in Lymph Nodes (OMLN) by Histological Review

Studies undertaken to detect occult lymph node metastases by routine histologic methods have generally been performed by cutting serial sections from all paraffin blocks containing lymph nodes, followed by routine staining and microscopic review (93). However, several studies have simply

rereviewed the original histologic slides. All of these studies have demonstrated that deposits of tumor can be detected using these methods. Between 7% and 33% of previously node-negative cases convert to node-positive after review. As reviewed by Neville (96), the mean conversion rate is approx 13%.

4.2. Detection of OMLN by IHC

Several investigators have used various antibodies in order to detect occult lymph node metastases in patients with breast cancer using IHC methods. In general, most studies have used antibodies specific for low molecular weight intermediate filament proteins to distinguish the epithelial tumor deposits from normal node elements. Other studies have used antimucin antibodies raised against human mammary carcinoma cells (100). In our own studies, we have used a cocktail of two anticytokeratin antibodies, AE1 and CAM5.2 (which in combination recognize CK-18 and CK-19, the predominant intermediate filament proteins in simple epithelial cells). While antibodies to cytokeratins have been reported to react with dendritic reticulum cells (with the possibility of producing false positive results) this has not been a significant problem in our own studies. Furthermore, as with the bone marrow examination previously described, the morphologic evaluation of the "positive" cells is critical; cells that do not possess the morphologic characteristics of malignant epithelial cells are not considered tumor cells in our studies.

Unlike routine histological evaluation for the detection of occult metastases in which multiple sections from each block are studied, most of the studies employing immunohistochemical techniques have tested only a single section. When a single section is studied, the percentage of patients who convert from node-negative to node-positive ranges from 14% to 30% with a mean conversion rate of 16% (82,96).

4.3. Detection of Occult Lymph Node Metastases by RT-PCR

Although it may at some point be possible to detect occult metastases in the lymph nodes of patients with breast cancer using PCR methods, this procedure continues to have major drawbacks compared with IHC. In addition to the problems of identifying appropriately sensitive and specific markers discussed above, the lymph nodes used must be fresh (or fresh frozen), and all lymph nodes in their entirety must be disaggregated to undergo RNA extraction (as we have shown that metastases in node-negative cases usually occurs to only one lymph node). Therefore, these lymph nodes will be unavailable for histologic review, and the method will test for both histologic and occult positives. This is in contrast to immunohistochemical techniques, which can be done on formalin-fixed, paraffinembedded tissue that has been routinely prepared for histologic evaluation.

4.4. Clinical Significance of OMLN in Patients with Breast Cancer

Although virtually all studies have demonstrated that lymph node metastases can be overlooked, there is a surprising disagreement about the prognostic importance of these occult tumor deposits. Many of the studies using routine histologic review of sections have found that the presence of OMLN does not influence the recurrence rates in a statistically significant way (84–86); several studies using immunohistochemical techniques have reported similar findings (90,91,98). To begin to understand this, a few basic observations need to be made. Many earlier studies have involved fewer than 100 patients; in fact some have involved even fewer than 50 patients. Cote and Groshen have demonstrated that even if the finding of occult lymph node metastases is prognostically important, there is no possibility that studies of the clinical impact of occult lymph node metastases involving few patients will provide statistically significant data (unpublished data). In fact, Fisher et al. (85) were the first to clearly point this out: "it has been mathematically estimated that differences in survival of 10%, if indeed they occur between the two groups (true lymph node negative versus occult lymph node positive), would require a study of approx 1400 cases." Therefore, studies involving a few patients are not suitable to address the issue of prognostic significance of occult lymph node metastases.

Investigators from the Ludwig Institute and International Breast Cancer study group have performed a definitive study of the importance of occult lymph node metastases in patients with node-negative breast cancer (87). They examined serial sections of 921 node-negative breast cancer patients by routine histological methods. Nine percent of these patients were found to have OMLN; these patients had a poorer disease-free (p = 0.003) and overall survival (p = 0.002) after 5 yr median follow-up, compared with patients whose nodes remained negative after serial sectioning. Six-year median follow-up data give even more conclusive evidence of the prognostic significance of occult lymph node metastases. Another large-scale study was performed (88). These investigators studied the lymph nodes from 1121 patients with primary operable breast cancer, by serial macroscopic sectioning; they found single OMLN in 120 patients. A significant difference in recurrence (p = 0.005) and survival (p = 0.04) was found between node-negative patients and those with single occult metastases. Immunohistochemical methods have also shown the prognostic significance of occult lymph node metastases (92,94,100). Trial V of the International (Ludwig) Breast Cancer Study for the presence of metastases showed that in lymph nodes from patients with breast cancer, occult metastases were associated with significantly poorer disease-free and overall survival in postmenopausal patients but not in premenopausal patients. Immunohistochemically detected OMLN remained an independent and highly

significant predictor of recurrence, even after control for tumor grade, tumor size, estrogen-receptor status, vascular invasion, and treatment (p = 0.007) (103).

It can be seen by these recent studies that the evidence is mounting that when sufficiently large patient populations are analyzed, OMLN is an independent indicator of the likelihood of disease progression. In addition, it appears that OMLN is a biologic event independent of bone marrow status and has an intermediate risk of recurrence associated with it (74).

4.4.1. SENTINEL LYMPH NODES

Axillary lymph node dissection (ALND) is the standard in care for patients with breast cancer. In more recent years the sentinel lymph node dissection (SLND) technique has been extensively studied. The technique uses isosulfan blue dye, a radiopharmaceutical, or a combination of both to locate and remove the first few lymph nodes that drain the tumor. These sentinel lymph nodes are considered the most likely to contain cancer cells. This procedure has the advantage of potentially saving patients with early stage breast cancer from having a complete axillary lymph node dissection and the subsequent morbidity that is associated with this procedure. Several studies have shown that this technique is highly accurate in staging patients. Giuliano et al. (104) showed a false-negative rate for patients who had an ALND performed after SLND of only 2%. Veronesi et al. (105) reported no false-negative sentinel lymph nodes among 45 patients with T1 breast cancer that measured less than 1.5 cm. The overall accuracy rate that included T3 tumors was 97%. In a more recent study they (106) examined 516 patients with breast cancer who had tumors that measured 2 cm or less in diameter. The overall accuracy of sentinel node status was 96.9%, the sensitivity was 91.2% and the specificity was 100%. They concluded that sentinel-node biopsy is an accurate screening of axillary node status in women with earlystage breast cancer. Furthermore, Alex and Krag (107) were able to identify the sentinel lymph nodes in 50 of the 70 patients they studied. Of these 50 patients 18 were found to have positive sentinel nodes that were predictive of the axillary status. Albertini et al. (108) were able to identify the sentinel lymph nodes in 57 of 62 (92%) patients. All patients with axillary nodal metastases were identified by the sentinel lymph node examination. However, a trial that looked at the sentinel lymph nodes from multiple institutions using only the radioisotope technique alone showed somewhat less promising results (109). In this study of 443 patients a sentinel lymph node was identified in 405 patients. These 405 then underwent a completion axillary lymph node dissection. Of these 405 there were 114 node-positive cases with 13 false-negative nodes for a false-negative rate of 11%.

In cases in which the routine histologic examination of the sentinel lymph node is negative, immunohistochemical evaluation for the presence of occult metastases may play an important role in the histopathologic evaluation of these patients. The reported rate of positivity for occult metastases in these histologically negative nodes ranges from 5% to 15% (92). It is clear that large-scale clinical trials are needed to determine the clinical relevance of these occult metastases positive sentinel lymph nodes.

4.4.2. Occult Metastases in Lymph Nodes in Patients with DCIS

The significance of finding occult metastases in patients with small tumors or comedo carcinoma is less clear than for patients with invasive carcinoma. The purpose for detecting occult metastases in lymph nodes, or in any other site for that matter, is to assess the risk for distant metastasis and death due to breast cancer. While most measures of DCIS outcome are focused on rates of local recurrence and the incidence of development of invasive disease, the critical issue with regard to lymph node analysis in DCIS is the distant outcome for the patient.

A number of studies have now addressed this issue. In an analysis from the NSABP Project B-17, it was found that for 814 patients with ductal carcinoma in situ followed for a median time of 7.5 yr, that the rate of distant metastasis was 1.8%, and deaths due to breast cancer was 1.8% (110). In another study 1002 women who were diagnosed with DCIS between 1986 and 1996 were followed for a median time of 4.25 yr. The rate of distant metastasis in this group was 2.4%, and deaths due to breast cancer was 1.5% (111). Finally, it should be noted that in all of these studies, a number of women subsequently developed invasive carcinoma and these women are included in the distant metastasis and death due to breast cancer rate. There is no question that some of the incidence of metastasis and death due to breast cancer is due to the subsequent development of either ipsilateral or contralateral invasive carcinoma. However, for the purposes of this evaluation, we have not separated out these patients, but have rather taken a "conservative" approach to the data analysis. Nevertheless, the data are virtually noncontroversial; systemic events in patients with DCIS are rare, under any circumstances. Therefore, detection of lymph node metastases, including occult metastases, does not appear to be an important procedure in patients diagnosed with DCIS overall.

However, all of the studies indicated above were performed in all patients with DCIS, without regard to subtype. Because the incidence of DCIS is increasing, and the proportion of women being diagnosed with breast cancer who have the diagnosis of pure DCIS is increasing as well, this subtype analysis may in fact become increasingly important. The question then becomes, do occult metastases occur in any patients with DCIS, and are there particular subtypes in which the risk for occult metastasis, and subsequent systemic events, is substantial?

There are preliminary data to support the idea that occult metastases may be detectable in at least in some types of DCIS (112,113). A positivity rate of up to 5% of nodal metastases in level 1 and 2 axillary dissections from patients with DCIS has been reported (114). However, because of the high morbidity associated with axillary lymph node dissections, and the low rate of recurrent disease in patients with DCIS, axillary lymph node dissection is not recommended as routine practice (115). The advent of the sentinel lymph node (SLN) biopsy technique has greatly reduced the morbidity associated with axillary lymph node sampling, offers a low-risk way of assessing lymph node status in DCIS. This may be of particular importance in patients that are more likely to have nodal involvement such as those with DCIS with an extensive intraductal component or comedo type (113,114). There is evidence that SLN biopsy complimented by IHC detection of occult metastases on these nodes may add to the relevance of the initial diagnosis and subsequent management of patients with DCIS (113,116). Studies to date show that a percentage of patients initially diagnosed with DCIS have tumor cells present in the sentinel nodes and that IHC evaluation of sentinel nodes is more sensitive than routine histology with serial sections at identifying these cases (113,116).

The prognostic significance of finding tumor cells in the sentinel nodes in patients with DCIS is not known. It is certain that not all positive cases will progress to more advanced disease; we know that in invasive cancer with overt nodal positivity, not all patients go on the develop recurrent disease. Nevertheless, large-scale prospective studies are needed to determine what clinical advantage, if any, exists in routinely analyzing sentinel lymph nodes for occult metastases in patients with DCIS.

5. OCCULT METASTASES IN THE BONE MARROW AND LYMPH NODES

While there is strong evidence to suggest that detection of either bone marrow or lymph node metastasis is clinically significant, it is possible that the detection of occult systemic (bone marrow) and regional (lymph node) metastases may provided complementary information in assessing the risk of an individual patient. Preliminary reports exist that support the hypothesis that occult metastases in the bone marrow and lymph nodes represent separate biologic pathways and are present in different subgroups of patients may have different prognostic significance. We studied 90 patients with node-negative breast cancer for the presence of both occult bone marrow and lymph node metastases (Table 2). OMLN were detected in 12 patients (13%) while OMBM were detected in 16 patients (18%). Both lymph node and bone marrow metastases were detected in only one patient (1%). Stratification showed that the presence of OMBM identified patients most likely

Lymph Wode vs Bone Warrow Occurt Wetastases							
OMLN/OMBM status	Patients	5-yr probability of recurring $(\pm SE)$	Relative risk of recurring	p-value			
OMLN-/OMBM-	62/90 (69%)	0.09 ± 0.04	1.00	< 0.00001			
OMLN ⁺	12/90 (13%)	0.31 ± 0.16	2.96				
OMBM ⁺	16 ^a /90 (18%)	0.50 ± 0.13	7.17				

Table 2Lymph Node vs Bone Marrow Occult Metastases

OMLN, Occult metastases in the lymph nodes; OMBM, occult metastases in the bone marrow; –, no occult metastases detected; +, occult metastases detected; SE, standard error. ^{*a*}Includes one OMLN⁺ patient.

to recur. Bone marrow negative patients could be stratified into risk groups based on their lymph node status (lymph node negative vs lymph node positive). Thus, three distinct groups were identified; those with detectable OMBM (high risk for recurrence), those with detectable occult metastases in the lymph nodes but none in the bone marrow (intermediate risk for recurrence), and those with no detectable occult metastases in either the lymph nodes or bone marrow (low risk for recurrence).

Gerber et al. (74) reported in their study that looked at 484 patients with pT1-2N0M0 breast cancer. Occult tumor cells were detected in 180 (37.2%) of the 484 patients; of these 126 (26%) patients had positive bone marrows only, 31 patients (6.4%) had positive lymph nodes only, and 23 patients (4.8%) had tumor cells found in both compartments. Patients who had occult metastases in and any compartment showed a decreased disease-free survival and the overall survival rates were found to be comparable top those patients who had one positive lymph node. Therefore, it is possible that occult metastatic tumor cells detected by simultaneous immunohistochemical analysis of axillary lymph nodes and bone marrow demonstrate independent metastatic pathways.

6. FUTURE DIRECTIONS

While the detection of occult metastases and the prognostic significance of their presence has been the primary focus to date, it is clear that this represents the tip of the iceberg in terms of the clinical applications possible. First, it is known that not all patients with overt metastases will suffer disease recurrence therefore, it is not reasonable to expect that all patients with occult metastases will recur. This is thought to occur because a proportion if the early-disseminated tumor cells are not viable or may be amenable to being destroyed by the patients immune system. Molecular markers that will predict which cells are likely to progress and which are not are currently being investigated. The relevance of erb-B2 overexpression on disseminated CK-18-positive breast cancer cells in the bone marrow of 52 patients with early-stage breast cancer using immunohistochemical double labeling to the p185erbB2 oncoprotein was examined (117). Expression of p185erb-B2 on cytokeratin-positive cells was detected in 60% of patients. It was determined that patients with p185erb-B2 positive bone marrow occult metastases developed fatal metastatic relapses more frequently than patients with p185erb-B2-negative bone marrow occult metastases. In addition, these markers may prove to be useful as specific targets to deliver chemotherapeutic agents (118). The analysis of occult metastatic cells opens a new avenue by which molecular determinants of both early tumor cell dissemination and subsequent outgrowth into overt metastases can be assessed. The identification of therapeutic targets such as HER2/neu MAGE-1 and E-Cadherin, for example, monitoring the elimination of bone metastases, and assessing treatment resistant tumor cell clones might help us to understand the current limitations of adjuvant systemic therapy.

The *MAGE-A* genes, genes that encode peptide antigens that can be recognized by autologous CTLs in the surface of the tumor cells in association with various classical HLA molecules, have been the subject of investigation (119). Because the *MAGE-A* genes are expressed in a variety of malignant tissues and absent in normal tissues other than the placenta and testis (120,121), their tumor-associated peptides could be used as targets for active immunotherapy. In addition, their expression analysis in malignancies could be of diagnostic and/or prognostic relevance. Preferential expression of *MAGE-A* has been observed in patients who are at a higher risk of recurrence: those harboring tumors with high levels of protease urokinase-type plasminogen activator and its inhibitor plasminogen activator inhibitor 1, high score of Ki-67 proliferation antigen, and a lesser degree if differentiation (119). These findings suggest a potential involvement of *MAGE-A* in tumor progression, with potential implications for active immunotherapy.

Another major concern, especially in terms of breast cancer, in which the disease is known to recur even after many years of remission even if adjuvant therapy was given, is that of tumor cell dormancy. Investigators at the Norwegian Radium Hospital (122) have examined the bone marrow aspirates from patients with breast cancer using IHC to detect cytokeratin positive cells before treatment, at the time of surgery and after treatment. At the time of primary diagnosis 22.5% of patients had OMBM, at the time of surgery 16.9% showed OMBM and after treatment 31.9% were positive. They found that 17.2% of patients who were initially negative for OMBM, became positive at the last aspiration, while 14.0% with an initially positive bone marrow results became negative (122). The theory behind this is that a population of tumor cells may exist in a "dormant" state, and these dormant

cells may not be responsive to traditional tumor cytotoxic therapies that act on proliferating or actively metabolizing cells. Thus, the study of dormancy in the early-disseminated tumor cell population may have important implications regarding the use (and specific type) of adjuvant therapy in patients with cancer. Dormancy is defined as the maintenance of a balance between proliferation and apoptosis, or a state of "quiescence." Dormant tumor cells may not be effectively treated by cytotoxic regimens that require actively metabolizing or proliferating cells. Studies are underway that will examine individual occult metastatic tumor cells detected for characteristics associated with tumor cell

According to some (123), the cytotoxic agents currently used for chemotherapy in high-risk breast cancer patients do not completely eliminate CK-positive tumor cells in the bone marrow. The presence of these tumor cells after chemotherapy is associated with poor prognosis. Thus bone marrow monitoring might help predict the response to systemic chemotherapy.

7. ONGOING PROSPECTIVE STUDIES

The true test of the prognostic importance of occult metastasis detection in both the bone marrow and lymph nodes will come from large-scale multiinstitutional studies. In the case of breast cancer, such a study is now well underway, under the auspices of the American College of Surgeons Oncology Group (ACOSOG), protocol Z0010 "A prognostic study of sentinel node and bone marrow micrometastases in women with clinical T1-2 N0 breast cancer." This study should provide definitive evidence of the prognostic significance of occult metastases in patients with breast cancer.

We have suggested a new concept in the staging of cancers using the TNM classification, where the traditional T (tumor), N (node), and M (metastasis) may be complemented by n and m (nodal and systemic occult metastases); that is TNnMm (28,124). With the results of larger studies on prognostic significance of occult metastases, either in bone marrow or lymph nodes, this staging may be applied clinically. The estimates of outcome for populations of patients may be narrowed down to those for subpopulations of patients (i.e., those with or without occult metastases).

8. PATIENT-SPECIFIC TREATMENT

The establishment of new techniques that can identify patients who are at risk of recurrence earlier coupled with the ability to determine the molecular profiles of the individual tumors may well enable patient specific treatment regimens to be developed. Treatment will be able to be tailored with regard to dormancy/proliferation state and biological markers of tumor progression that may be present. In addition, the development of therapeutic agents designed to be delivered specifically to the tumor cells will enable more effective elimination of the tumor cells while decreasing the morbidity associated with conventional chemotherapeutic agents that target proliferating cells. Thus more effective and less toxic adjuvant therapy may be provided for women with primary breast cancer.

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VI FUTURE DIRECTIONS

16 New Technologies/ New Markers/ New Challenges

Daniel F. Hayes, MD and Giampetro Gasparini, MD

CONTENTS

INTRODUCTION FUTURE DIRECTIONS CONCLUSIONS REFERENCES

Key Words: Biomarkers; technologies; development.

1. INTRODUCTION

The field of tumor markers is enigmatic. On the one hand, the explosion of technology in medicine has provided an enormous number of complex tools that might be used to screen for, diagnose, or prevent a newly diagnosed cancer and/or its metastases. Indeed, the preceding chapters have highlighted remarkable progress in these technologies, and potential areas in which they might be useful. On the other hand, very few markers have actually been accepted for routine clinical use in breast cancer. The American Society of Clinical Oncology Tumor Marker Expert Guidelines Panel has issued three sets of guidelines since its inception in 1996. In the initial set of guidelines, the use of estrogen and progesterone receptors (ERs, PgRs) to select patients for hormone therapy and the use of circulating CA15-3 and/or carcinoembryonic antigen (CEA) to monitor selected patients with

Cancer Drug Discovery and Development: Biomarkers in Breast Cancer: Molecular Diagnostics for Predicting and Monitoring Therapeutic Effect Edited by: G. Gasparini and D. F. Hayes © Humana Press Inc., Totowa, NJ metastatic disease were the only recommended markers (1). In subsequent updates, the Panel recommended routine measurement of *HER2* in cancer tissue to select trastuzumab for patients with metastatic disease (2,3). None of the other markers that have been reported and proposed for breast cancer was felt to be sufficiently validated for routine use by the Panel. Similar recommendations have been made by the College of American Pathologists (4).

Why are these guidelines so conservative? There are several answers to this question, and each of these questions should serve as the basis for future studies (5).

1. A marker is helpful only if it separates an entire population into two different groups whose outcome is likely to be so different that one group might be treated differently than another. Both ER and HER2 are classic examples of this category. Patients with ER-negative tumors appear very unlikely to benefit from hormone therapy (6), and, likewise, it appears that patients with HER2-low or -negative cancers are very unlikely to benefit from trastuzumab (5,7).

Therefore, future studies should be focused on identifying markers with sufficient strength that outcomes of patients who are "positive" (by whatever criteria used) have sufficiently different outcomes that a clinician would treat them differently than those who are "negative." It is certainly possible that selected, single-gene or single-protein markers might fall into this category. For example, data from Europe support the notion that patients whose tumors do not express urokinase plasminogen activator (uPA) and the plasminogen activator inhibitor 1 (PAI-1) proteins have a sufficiently favorable prognosis that the benefits from chemotherapy would be so low that they might forego this toxic form of therapy (8,9).

The new technologies that permit high-throughput analysis may well identify such a marker. These technologies have two promises: new gene or protein identification and pattern recognition. Currently, the latter has achieved the most notoriety. High-profile publications and presentations have suggested that selected combinations of multiplex gene expressions are associated with very good or poor prognosis (10-12). However, although promising, these observations require validation in prospective, properly designed clinical trials.

Proteomics also offers great promise in identifying specific patterns that might be helpful in the clinic (13, 14). However, this field is in its infancy, with many methods and techniques, and at this writing there have been few if any studies that suggest any clinical utility for proteomic pattern recognition in breast cancer.

The use of high-throughput technologies also provides the opportunity to identify new, previously unrecognized genes and their products that might be of value. However, so far, none has been recognized that appear to be of clinical value.

2. The assay must be reliable and reproducible. With few exceptions, tumor markers seem to pass through a "life cycle" in which the original report is extraordinarily positive with great acclaim, but subsequent studies fail to live up to the promise. There are fundamentally three reasons for this conundrum: (1) technical variability of the assay; (2) variations in the manner in which different assays for the same marker are performed; and (3) inadequate and differences in study design. These effects of these concerns are often underestimated in the excitement of apparently "positive" results, but ultimately they are paramount in the consideration of whether a new marker is or is not helpful in caring for patients. Thus, as new techniques and assays are developed, it is important to demonstrate that not only are the result promising, but that they are reproducible. The first of these three requirements, technical variability, is perhaps the most easily accomplished, although even this issue is not always demonstrated. The latter two issues are more common, and perhaps more important, reasons that new markers are so slowly adopted. Because of competition among scientists and commercial interests, different assays are often developed to evaluate the same marker. For example, HER2 status can be determined by examination of breast cancer tissue amplification of the erb-B2 gene using a variety of techniques including Southern blotting, slot-blot quantification, or fluorescence in situ hybridization, and by evaluation of the protein using Western blotting, immunohistochemistry, immunofluorescence, or enzymelinked immunosorbent assays (ELISA). Moreover, circulating extracellular domain of HER2 can be quantified in human serum using ELISA (15). Although they are all correlated, each of these assays, which in one way or another provides an indication of overproduction of HER2, appears to differ from the other and to provide different results in regards to prediction of outcome. Thus, it is not surprising that results from study to study are not validated, if the assays that are being compared are not identical.

Another technical issue that hinders reproducibility is related to selection of the cutoff that distinguishes positive from negative populations. Several means of doing so are employed. One method is to arbitrarily select a cutoff, based on some preconceived reason, such as the mean level of the assay in an affected population or the mean plus two standard deviations of the level in an unaffected population. A second method is to test several potential cutoffs in one population, selecting the one that appears most robust in regard to separation of the outcomes of the two groups or in regards to apparent statistical significance. Regardless of the method used, it is essential to validate the results in a separate group of patients.

Finally, the results are most likely to be valid if they are studied in the context of a plausible hypothesis that is prospectively addressed. Many published studies report results related to hypotheses that are retrospectively derived from the observed data. Although such studies are valuable

to generate hypotheses, these observations must be prospectively validated in subsequent, well-designed studies (5,16). Unfortunately, most tumor marker studies are performed using archived specimens collected for reasons unrelated to the study under question. Therefore it is difficult to validate observations, requiring and time-consuming prospective studies. Nonetheless, failure to do so often leaves the marker hanging, with the original exciting results, generated from the retrospective exploratory study, lacking validation.

2. FUTURE DIRECTIONS

The previous caveats notwithstanding, technological advances do offer great promise for individualization of therapy for breast cancer. We believe the following areas of research are particularly exciting.

2.1. Genomics/Proteomics

The ability to examine abnormalities in multiple genes and/or their products simultaneously has permitted avenues of discovery that are unprecedented. Clearly, multiplex gene amplification and gene expression, studying tens, hundreds, and even thousands of genes and/or proteins simultaneously have the potential to move the field exponentially when compared to a gene-by-gene approach. These methods lead to two possible advances: Discovery of new, single genes that may play an important role in breast cancer; and identification of patterns of genes that, even if their individual roles and contributions are uncertain, can act as a single tumor marker when considered in aggregate. The former approach offers the promise not only of improving diagnostics, but of providing clues to potential targets for therapeutic research. The latter, if reproducible and validated, may become the sine que non for powerful markers that clearly separate populations with favorable from unfavorable outcomes to the extent that they would be treated much differently. Most of the currently available and promising data have come from genomic studies, either of frozen (10,11,17) or fixed tissue (12). However, studies of multiple proteins, including evaluation of posttranslational modification, may well be a much more powerful and accurate strategy in the long run, although the methodologies of this approach are still in their infancies.

2.2. Minimal Detectable Disease

Identification of apparently malignant cells in the circulation or bone marrow has been possible for >130 yr (18). Recent technologies have substantially improved the accuracy and reliability of enumerating such cells, using immunoseparation techniques or reverse transcriptase-polymerase chain reaction (RT-PCR) (19–21). Perhaps even more exciting is the ability

to genotype and phenotype these cells once they are identified (22-25). Although clinical utility of this technology has not been shown, the ability to perform "real time" evaluation of the current biology (e.g., *HER2* status or the level of apoptosis) of a patient's cancer is of great promise.

2.3. Pharmacogenomics

Nearly all of this volume has addressed somatic changes within the cancer itself that might predict its natural or treated history. Increasingly, technological advances in genomics and proteomics have permitted studies of inherited, germ-line polymorphisms that may affect response to or toxicity from therapy. These single-nucleotide polymorphisms (SNPs) result in subtle but often important amino acid substitutions within genes that are important for activation and/or metabolism of the drug, or within genes that may serve as the target for the drug (26). Only preliminary studies of these pharmacogenomic (or pharmacogenetic) changes have been reported, both for chemotherapy and for endocrine treatments, such as tamoxifen (27,28).

3. CONCLUSIONS

In summary, incredible technological advances have opened the door for a potential sea change of evaluation and treatment of patients with breast cancer. These advances should help to improve the ability to evaluate risk, screen for new cancers, estimate prognosis and prediction of response to therapy, monitor patients during therapy, and even serve as targets for novel treatments. However, for any of these to take its place in clinical medicine, it will need to be carefully studied in rigorously designed clinical trials based on preexisting hypotheses, with sophisticated statistical analyses, taking into account other factors and treatment effects.

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Index

A

Amphiregulin, breast cancer expression, 180 Angiogenesis biomarker, see Vascular endothelial growth factor inhibitors endostatin, 284, 285 paclitaxel, 283 thalidomide, 284 vascular endothelial growth factor trials monoclonal antibody targeting, 279, 283, 284 overview, 277, 278 tyrosine kinase inhibitors, 279-281 neovascularization process in cancer, 268, 269 regulators, 268, 269 tumor growth and progression role, 270, 271 Anthracycline-based regimens, HER2/neu response studies cyclophosphamide, methotrexate, 5-fluorouracil regimen comparison, 143-146 dose response, 146, 147 overview, 143

B

Biomarker, see also specific biomarkers clinical trials caveats, 5 design, *see* Clinical trial design levels of evidence, 29 criteria for useful biomarkers assay reliability and reproducibility, 327, 328 group outcomes, 326, 327 definition, 4 immunoassay, *see* Immunoassay, biomarkers Bone marrow tumor cells, *see* Disseminated tumor cells

С

CA 15-3, early breast cancer recurrence detection, 258, 259 Carcinoembryonic antigen (CEA), early breast cancer recurrence detection, 258, 259 CBCTR, see Cooperative Breast Cancer **Tissue Resource** CEA, see Carcinoembryonic antigen Cell proliferation assays antigen-related markers, 86, 87 flow cytometry of nuclear DNA content and S-phase fraction, 85, 86 M phase, 84 overview, 82-84 proliferation index comparisons clinical utility, 100-102 correlations, 88, 89 prognostic significance, 91–97 reproducibility and quality control, 87, 88

risk profiles, 89, 91 treatment response, 97-100 S phase, 85, 86 cell cycle phases, 82, 84 *c-erb*-B2, see HER2/neu CGH, see Comparative genomic hybridization Chromogenic in situ hybridization (CISH), Her2 gene amplification analysis, 72, 74 Circulating tumor cells CK-19 marker detection with reverse transcriptasepolymerase chain reaction, 221 detection flow cytometry, 295 immunohistochemistry, 295, 296, 298, 299 immunomagnetic separation, 299, 300 reverse transcriptase-polymerase chain reaction, 297, 298, 299 immunocytochemistry, 217, 220 patient-specific therapy design, 314 prognostic value, 300, 301 prospects for study, 224-227, 311-313 selection strategies, 220, 221 CISH, see Chromogenic in situ hybridization CK-19, occult metastatic cell marker detection with reverse transcriptase-polymerase chain reaction, 221, 297, 298 Clinical trial design biomarker caveats, 5 biomarker validation studies examples of patient identification node-negative patients not requiring chemotherapy, 12, 13 node-positive patients benefiting from specific regimens, 14, 15

node-positive patients not benefiting from chemotherapy, 13, 14 guidelines, 10-12, 19 phase III trials, 10, 11 retrospective study analysis of biomarkers, 6-10 CMF, see Cyclophosphamide, methotrexate, 5-fluorouracil *c-neu, see* HER2/neu Comparative genomic hybridization (CGH), Her2 gene amplification analysis, 75, 76 Cooperative Breast Cancer Tissue Resource (CBCTR) array characteristics, 40, 43 origins, 39 Cross-validation, leave-one-out cross-validation, 8, 9 Cyclophosphamide, methotrexate, 5fluorouracil (CMF) circulating Her2/neu extracellular domain as prediction indicator, 248 HER2/neu response studies, 141–143

D

DCIS, see Ductal carcinoma in situ Disseminated tumor cells (DTCs) bone marrow tumor cell clinical significance, 214, 215, 222-224, 226, 227, 301-304, 311 CK-19 marker detection with reverse transcriptasepolymerase chain reaction, 221, 297, 298 detection in bone marrow flow cytometry, 295 immunohistochemistry, 295, 296, 301 reverse transcriptasepolymerase chain reaction, 297, 298 tumor burden effects, 304, 305

histopathology, 215 immunocytochemistry, 215-217 patient-specific therapy design, 314 prospective studies, 313 prospects for study, 224-227, 311-313 selection strategies, 220, 221 DNA methylation, metastatic breast cancer, 222 DNA microarray advantages, 58 biomarker discovery, 5, 6 breast tumor analysis classification, 54, 55 historical perspective, 52 pathogenesis and progression studies, 52-54 prognosis and prediction, 55, 56 combinatorial analysis, 56, 57 data analysis, 51 DNA chips, 49, 50 messenger RNA correlation with protein levels, 57 microdissected samples, 57, 58 pooling of samples, 58 principles, 48, 49 prospects, 59 reference samples, 58 sample preparation, 50 Docetaxel circulating Her2/neu extracellular domain as prediction indicator, 249-251 HER2/neu response studies, 147–149 DTCs, see Disseminated tumor cells Ductal carcinoma in situ (DCIS) epidermal growth factor receptor expression, 183 lymph node occult metastasis, 309-311

E

EGF, *see* Epidermal growth factor EGFR, *see* Epidermal growth factor receptor ELISA, see Enzyme-linked immunosorbent assay Endostatin, angiogenesis inhibition, 284, 285 Enzyme-linked immunosorbent assay (ELISA) Her2/neu extracellular domain, 239,240 plasminogen activator inhibitor-1, 120, 121 principles, 19, 20 urokinase plasminogen activator, 120, 121 Epidermal growth factor (EGF) breast cancer expression, 179 structure, 178 transforming growth factor- α homology, 179 Epidermal growth factor receptor (EGFR) breast cancer expression, 181, 182 breast development role, 182 drug resistance role, 184 ductal carcinoma in situ expression, 183 EGFR-2, see HER2/neu estrogen effects on expression, 185 family members, 181 ligands, 179–181 signaling, 160, 181, 182, 184 tyrosine kinase inhibitor targeting, 183 erb-B2, see HER2/neu

F

FISH, see Fluorescence in situ hybridization Flow cytometry nuclear DNA content and S-phase fraction overview, 85, 86 proliferation index comparisons, 87–89, 91–93, 97–102 occult metastasis detection, 295 Fluorescence *in situ* hybridization (FISH), *Her2* gene amplification analysis advantages, 68–70 limitations, 72, 74 optimization for archived specimens, 72, 73 reproducibility, 71, 72 trastuzumab response studies, 169, 170, 173

G

Gene amplification breast cancer amplified genes, 66 Her2 amplification amplicon gene types, 67 chromogenic in situ hybridization, 72,74 clinical significance, 67 comparative genomic hybridization, 75, 76 fluorescence in situ hybridization advantages, 68-70 limitations, 72, 74 optimization for archived specimens, 72, 73 reproducibility, 71, 72 immunohistochemistry, 68-70 metastasis samples, 67, 68 real-time reverse transcriptase polymerase chain reaction, 74,75 trastuzumab response monitoring, 68-70 Gene expression profiling, see DNA microarray Genomics, breast cancer treatment prospects, 328

Η

HER2/neu activation and signaling, 160, 237, 238 circulating extracellular domain assays, 239, 240 early recurrence detection, 258, 259 prevalence in breast cancer, 240-242 prognostic value, 242–245 prospects for study, 259 shedding, 238 treatment response prediction chemotherapy, 248-251 hormone modulation therapy, 246, 247 metastatic breast cancer, 252, 256-258 trastuzumab therapy, 251, 252 estrogen effects on expression, 185 function, 130 gene amplification amplicon gene types, 67 chromogenic in situ hybridization, 72,74 clinical significance, 67 comparative genomic hybridization, 75, 76 fluorescence in situ hybridization advantages, 68-70 limitations, 72, 74 optimization for archived specimens, 72, 73 reproducibility, 71, 72 immunohistochemistry, 68-70 metastasis samples, 67, 68 real-time reverse transcriptase polymerase chain reaction, 74,75 trastuzumab response monitoring, 68-70 gene locus, 236 structure, 130, 160, 236, 237 therapeutic targeting in conjunction with epidermal growth factor receptor, 185, 186

therapeutic targeting, see Trastuzumab treatment response monitoring in breast cancer chemotherapy response anthracycline-based regimens, 143–147 clinical application, 149, 150 cyclophosphamide, methotrexate, 5-fluorouracil therapy, 141 - 143preclinical data, 141 taxane-based regimens, 147 - 149clinical recommendations, 151, 152 controversies, 130, 131 endocrine manipulation response adjuvant chemotherapy studies, 138-140 drugs, 132, 133 metastatic breast cancer, 133-135, 138 preclinical data, 133 study limitations, 140 level of evidence studies, 131 study design, 131, 132 trastuzumab response, 150, 151 Herceptin[®], see Trastuzumab Heregulins, breast cancer expression, 180, 181

I

IHC, *see* Immunohistochemistry Immunoassay, biomarkers accuracy definition, 23 high-dose hook effect, 24 interferences, 24 linearity, 23 recovery, 23, 24 ideal criteria, 18

interpretation, 26 minimal consistency criteria, 26 normalization of results, 28 platforms, 19, 20 precision, 25, 26 quality control daily performance monitoring, 27,28 external quality assessment, 28, 29 external quality control, 27, 28 internal quality control, 27 overview, 26, 27 samples, 27 sampling bias and tissue processing, 21,22 sensitivity, 25 specificity, 24, 25 standards and calibration, 22, 23 variability of results, 20, 21 Immunohistochemistry (IHC) HER2/neu antigen expression, 162, 163, 168, 169 gene amplification analysis, 68-70 occult metastasis detection bone marrow tumor cells, 215-217, 295, 296, 301 lymph node metastasis, 295, 296, 306, 307 peripheral blood, 217, 220, 295, 296, 298, 299 p53 mutation effects on expression, 195 standardization, 195, 196 Immunomagnetic separation, occult metastasis detection in peripheral

K

Ki-67 proliferating cell marker, 86, 87

blood, 299, 300

L

Leave-one-out cross-validation (LOOCV), accuracy prediction, 8,9 Letrozole, circulating Her2/neu extracellular domain as prediction indicator, 246, 247 LOOCV, see Leave-one-out crossvalidation Lymph node metastasis clinical significance in breast cancer, 214, 294, 307-311 detection flow cytometry, 295 histological review, 305, 306 immunohistochemistry, 295, 296, 306, 307 limitations, 305 reverse transcriptasepolymerase chain reaction, 297, 298, 307 ductal carcinoma in situ, 309-311 patient-specific therapy design, 314 prospective studies, 313 prospects for study, 311-313 sentinel lymph nodes, 308, 309

M

MAI, see Mitotic activity index Metastasis circulating Her2/neu extracellular domain as treatment prediction indicator Herceptin, 257, 258 hormone or chemotherapy, 252, 256 DNA methylation, 222 Her2/neu gene amplification, 67, 68 endocrine manipulation response studies, 133–135, 138 occult metastatic cells, *see* Circulating tumor cells; Disseminated tumor cells; Lymph node metastasis vascular endothelial growth factor expression, 271 Microarray, *see* DNA microarray; Tissue microarray Mitotic activity index (MAI) determination, 84 proliferation index comparisons, 87–89, 91–93, 97–102

Ν

National Surgical Adjuvant Breast and Bowel Project (NSABP), tissue array resources, 43 Neu, *see* HER2/neu NSABP, see National Surgical Adjuvant Breast and Bowel Project

0

Occult metastasis, *see* Circulating tumor cells; Disseminated tumor cells; Lymph node metastasis

P

p53 breast cancer gene mutations overexpression correlation, 195 sequencing, 196, 197 types and clinical relevance, 197, 198 prognostic value studies, 194, 204, 205 prospects for study, 204, 206 treatment response studies biological rationale, 198, 201 limitations, 194

p53-independent drugs, 201 study design, 202, 203 functions, 194, 195, 204 immunohistochemistry mutation effects on expression, 195 standardization, 195, 196 single-stranded conformational polymorphism studies, 196 Paclitaxel angiogenesis inhibition, 283 circulating Her2/neu extracellular domain as prediction indicator, 249-251 HER2/neu response studies, 147–149 p53 gene mutation effects, 201 PAI-1, see Plasminogen activator inhibitor-1 Peripheral blood tumor cells, see Circulating tumor cells Pharmacogenomics, breast cancer treatment prospects, 328 Plasminogen activator inhibitor-1 (PAI-1) breast cancer expression cell types and distribution in tumors, 114, 115 modulators, 114, 115 prognostic value, 118–120 treatment response prediction, 121, 122 tumor grade effects, 114 enzyme-linked immunosorbent assay, 120, 121 functions, 113, 114 protein-protein interactions, 113 structure, 113 Polymerase chain reaction, see Reverse transcriptase-polymerase chain reaction Proliferative activity, see Cell proliferation

Proteomics, breast cancer treatment prospects, 328

Q

QC, see Quality control Quality control (QC) immunoassays for biomarkers daily performance monitoring, 27, 28 external quality assessment, 28, 29 external quality control, 27, 28 internal quality control, 27, 28 internal quality control, 27 overview, 26, 27 samples, 27 proliferation indices, 87, 88

R

Reverse transcriptase-polymerase chain reaction (RT-PCR) *Her2* gene amplification analysis with real-time reverse transcriptase polymerase chain reaction, 74, 75 occult metastatic cell marker detection with reverse transcriptase-polymerase chain reaction bone marrow tumor cells, 297, 298 lymph node metastasis, 297, 298, 307 overview, 221 peripheral blood tumor cells, 297, 298, 299 RT-PCR, see Reverse transcriptasepolymerase chain reaction

S

Single-stranded conformational polymorphism (SSCP), p53 studies, 196

Т

Tamoxifen circulating Her2/neu extracellular domain as prediction indicator, 246, 247 HER2/neu response studies adjuvant chemotherapy studies, 138 - 140metastatic breast cancer, 133–135, 138 preclinical data, 133 study limitations, 140 transforming growth factor- α inhibition, 179 Taxanes, see Docetaxel; Paclitaxel TGF-α, see Transforming growth factor-α Thalidomide, angiogenesis inhibition, 284 Thymidine labeling index (TLI) determination, 85 proliferation index comparisons, 87-89, 91-93, 97-102 Tissue microarray (TMA) advantages, 32 analysis systems, 37, 38 applications amplicon survey, 39 clinical laboratory, 37, 39 approaches, 33, 34 array sources commercial sources, 40-42 Cooperative Breast Cancer Tissue Resource, 39, 40, 43 National Surgical Adjuvant Breast and Bowel Project, 43 arrayers Advanced Tissue Arrayer, 34

Beecher Instruments Automated Arrayer ATA-27, 34,35 manual arrayer, 34 core numbers per case, 35 donor blocks standard operating procedure for array construction, 35, 36 submission refusal in multicenter trials, 36, 37 historical perspective, 32, 33 limitations, 39 prospects, 39, 40 TKIs, see Tyrosine kinase inhibitors TLI, see Thymidine labeling index TMA, see Tissue microarray Transforming growth factor- α (TGF- α) breast cancer expression, 179, 180 epidermal growth factor homology, 179 tamoxifen inhibition, 179 Trastuzumab circulating Her2/neu extracellular domain as prediction indicator, 251, 252 development antibody screening, 161, 162 antigen, 161 clinical trials, 163-168 humanization, 164 immunohistochemistry assay for antigen expression, 162, 163, 168, 169 fluorescence in situ hybridization studies of response, 169, 170, 173 Her2 gene amplification studies, 68 - 70HER2/neu response studies, 150, 151 history of development, 161 mechanism of action, 67, 68, 183, 238 prospects for study, 173

Tyrosine kinase inhibitors (TKIs) epidermal growth factor receptor targeting, 183 vascular endothelial growth factor receptor targeting, 279–281

U

uPA, see Urokinase plasminogen activator Urokinase plasminogen activator (uPA) breast cancer expression cell types and distribution in tumors, 114, 115 modulators, 114, 115 prognostic value, 116–118 treatment response prediction, 121, 122 tumor grade effects, 114 enzyme-linked immunosorbent assay, 120, 121 function, 112, 113 inhibitor, see Plasminogen activator inhibitor-1 metastasis role, 112 receptor binding, 113 structure, 112 therapeutic targeting, 123

angiogenesis role, 268–270 cancer biomarker validation, 271, 272 circulating biomarker studies endostatin level correlation, 282 individual variability, 277 inhibition trials monoclonal antibody targeting, 279, 283, 284 overview, 277, 278 tyrosine kinase inhibitors, 279 - 281measurement variability, 273, 277 prognostic-predictive value, 281-285 prospects for study, 285, 286 sample collection, 274–276 serum versus plasma levels, 281, 282 expression regulation, 271 isoforms, 270, 271 metastasis expression, 271 receptors, 270, 271 VEGF, see Vascular endothelial growth factor

W

Western blot, Her2/neu extracellular domain, 239

v

Vascular endothelial growth factor (VEGF)