Gene expression profiles do not consistently predict the clinical treatment response in locally advanced breast cancer

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Abstract
Neoadjuvant treatment offers an opportunity to correlate molecular variables to treatment response and to explore mechanisms of drug resistance in vivo. Here, we present a statistical analysis of large-scale gene expression patterns and their relationship to response following neoadjuvant chemotherapy in locally advanced breast cancers. We analyzed cDNA expression data from 81 tumors from two patient series, one treated with doxorubicin alone (51) and the other treated with 5-fluorouracil and mitomycin (30), and both were previously studied for correlations between response to chemotherapy and tumor subtype. Using supervised analysis, we could not uncover a gene profile among patients with luminal B type tumors treated with doxorubicin (5 of 8 patients; \( P = 0.0078 \); however, aside from these two observations, no other consistent associations between response to chemotherapy and tumor subtype were observed. These specific associations could possibly be explained by covariance with TP53 mutation status, which also correlated with tumor subtype. Using supervised analysis, we could not uncover a gene profile that could reliably (> 70% accuracy and specificity) predict response to either treatment regimen. [Mol Cancer Ther 2006;5(11):2914–8]

Introduction
Resistance to cytotoxic compounds is a main reason for therapy failure in most malignancies, including breast cancer. In vitro experiments as well as studies in animal models have shown that mutations in the TP53 gene are associated with chemoresistance (1). Molecular studies of tumors from patients treated with neoadjuvant chemotherapy using either doxorubicin monotherapy or 5-fluorouracil and mitomycin (FUMI) in concert revealed that TP53 mutations affecting the DNA-binding domain of the protein correlate with drug resistance (2–4). However, neither in these tumors nor in the studies reported by others (5) did mutations in TP53 unequivocally predict drug resistance, suggesting that other interactions and genes must be involved (6).

By subjecting the same tumors characterized for TP53 mutations in relation to chemotherapy response to DNA microarray analysis, we were able to classify tumors into five distinct subtypes based on their gene expression patterns (7). This classification showed prognostic effect with respect to relapse-free as well as overall survival in our cohort (8) and also in series of patients examined by other investigators (9). The prognostic significance of gene expression profiles has been well documented with respect to breast cancer (10–13) as well as other malignancies (14–17). Although these findings confirm the biological relevance of such genomic analyses, a prognostic factor provides no specific information about responsiveness to specific treatments and should be distinguished from a “predictive factor” (18, 19). Knowledge about the value of genome-wide expression analyses in predicting treatment response in breast cancer has resulted in at least to two studies correlating gene expression profiles with sensitivity to taxane monotherapy (20, 21) and three studies (22–24) reporting sensitivity to anthracycline combination regimens containing either cyclophosphamide or a taxane. However, the predictive powers achieved in any of these studies do not allow clinical implementation without further evaluations.
The aim of this study was to examine the potential of gene expression profiles as predictive factors of drug sensitivity in two uniformly treated breast cancer cohorts previously characterized for the predictive value of TP53 mutations and for the prognostic importance of gene expression profiles. Similar to findings by others, we found gene expression profiles defined by response-guided supervised analysis to be limited with respect to predicting therapy response.

Materials and Methods

Patient and Treatment Information

The patients included in this study were part of two prospective studies evaluating predictive factors for response to chemotherapy in locally advanced breast cancer (T3/T4 and/or N2). From one (doxorubicin series), we analyzed tumor samples from a subgroup consisting of 51 patients who were treated with doxorubicin monotherapy weekly in the neoadjuvant setting, scheduled for 16 weeks with 4 weekly assessments of clinical response (3). From the second similar study (FUMI series), we analyzed tumors from 30 patients who were treated with FUMI at 3-week intervals (4). Because these protocols were applied before implementation of the “Response Evaluation Criteria in Solid Tumors” criteria (25), for both studies, the response rates were classified according to the International Union Against Cancer criteria (26). Thus, responses were classified as partial response (reduction >50% in the sum of all tumor lesions, calculated for each as the product of the largest diameter and the one perpendicular to it), progressive disease (increase in the diameter product of any individual tumor lesion by >25%), or stable disease (anything between partial response and progressive disease). Therapy was terminated immediately in case progressive disease was revealed. An overview of patient characteristics is shown in Table 1, and a complete listing of all tumors and experiments is available in Supplementary Table S1.8

Microarray Analysis

Gene expression data were collected using cDNA arrays produced at the Stanford Functional Genomics Facility.9 The procedures used, including RNA extraction, hybridization, and data processing, have been described previously (7, 8) and are available at the Stanford Genomics Breast Cancer Consortium Portal Web site.10 The common set of genes used for the doxorubicin series totaled ~8,000, whereas for the FUMI series this number was ~30,000 due to more recent production lots of cDNA microarrays. Specifically, for these analyses, the background-subtracted, lowess-normalized (27) log2 ratio (Cy5/Cy3) intensity values were first filtered to select genes that had a signal intensity of at least 30 units above background in both channels. Only genes that met these criteria in at least 70% of the total data set were included for subsequent analysis, which totaled 4,424 probes for the entire data set. Next, missing values were imputed using the k-nearest neighbor imputation algorithm (28). Gene annotation from each data set was translated to UniGene Cluster IDs using the SOURCE database (29). Multiple occurrences of a UniGene Cluster IDs were collapsed by the median value for that ID within an experiment set.

Statistical Analysis

Relationships between gene expression profiles and response to chemotherapy were analyzed using “nearest shrunken centroid classifier” [prediction analysis for microarrays (PAM); ref. 30]. In addition, several other supervised prediction methods were used: recursive sample classification and gene selection with SVM for microarray data (r-SVM; ref. 31), Random Forest by Salford Systems (32), a k-nearest neighbor classifier with either Euclidean distance or one-minus-Spearman correlation as the distance function, and a class nearest centroid metric with either Euclidean distance or one-minus-Spearman correlation as the distance function (See Supplementary data for a more detailed description of the various methods; ref. 33).8 As discussed elsewhere (19), the terms partial response and stable disease are pragmatic terms that describe a status of tumor “growth arrest” with or without a certain degree of macroscopic reduction in tumor size; the discrimination between the two may often be arbitrary. However, progressive disease tumors are distinctive and easily discriminated from the other groups; therefore, our primary statistical analyses aimed at comparing progressive disease tumors versus the others. Finally, because these experiments were done across many different production lots of microarrays, we attempted to correct for systematic array batch bias by using “distance-weighted discrimination” (34).

Table 1. Clinical characteristics of patients included in this study

<table>
<thead>
<tr>
<th>Response</th>
<th>Doxorubicin (51)</th>
<th>FUMI (30)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PD</td>
<td>8</td>
<td>7</td>
</tr>
<tr>
<td>SD</td>
<td>22</td>
<td>13</td>
</tr>
<tr>
<td>PR</td>
<td>21</td>
<td>10</td>
</tr>
<tr>
<td>Histology</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ductal</td>
<td>46</td>
<td>26</td>
</tr>
<tr>
<td>Other</td>
<td>5</td>
<td>4</td>
</tr>
<tr>
<td>TP53 mutations</td>
<td>24</td>
<td>16</td>
</tr>
<tr>
<td>Median age</td>
<td>62 (32–85)</td>
<td>64 (37–82)</td>
</tr>
<tr>
<td>Median OS (mo)</td>
<td>33 (7–92)</td>
<td>24 (3–54)</td>
</tr>
<tr>
<td>No. relapses</td>
<td>26</td>
<td>17</td>
</tr>
</tbody>
</table>

Abbreviations: PD, progressive disease; SD, stable disease; PR, partial response; OS, overall survival.
Results and Discussion

Prognostic versus Predictive Factors

Several gene expression-based classification schemes for various cancer types have emerged from DNA microarray studies over the last years. The biological importance of these classifications is highlighted by two significant observations: first, the possibility that, for each cancer type, including breast (9, 12, 35, 36), lung (15), lymphoma (14, 17), and head and neck (33), it is possible to classify individual tumors into groups characterized by distinct gene profiles, and second, the fact that these classifications provide prognostic information. These analyses, however, have thus far been of limited value for predicting therapeutic response in individual patients. A prognostic factor is traditionally associated with disease-free or overall survival (in the absence of systemic adjuvant therapy), whereas a predictive factor predicts response, or lack of, to a particular treatment (19). Although numerous prognostic factors have been identified in breast cancer, no predictive factors have been generally accepted thus far, with the exception of estrogen receptor-α and progesterone receptor for endocrine therapy and HER2 for trastuzumab. In previous studies (3, 4), we found mutations in the TP53 gene affecting the L2/L3 domains of the protein to be associated with nonresponse to treatment with doxorubicin or FUMI. However, such mutations were only predictive for nonresponse in 60% of the progressive disease tumors. Although our findings strongly advocate a role of the p53 pathway in response to these therapies in breast cancer, they also suggest that other genes must also be involved (6).

Response to Therapy across Different Molecular Subtypes

In this study, we conducted statistical analyses of gene expression data from altogether 81 tumors, which represents one of the largest studies to explore the predictive value of gene expression profiles in breast cancer (23, 37). Response to therapy in tumors across the different, previously defined tumor subgroups is depicted in Fig. 1. One of 25 tumors belonging to the luminal A subgroup versus 7 of 26 tumors in all other groups were non-responding (progressive disease) to doxorubicin ($P = 0.0496$, two-sided Fisher’s exact test); for the patients

![Figure 1](https://example.com/figure1.png)

**Figure 1.** Distribution of progressive disease (PD), stable disease (SD), and partial response (PR) tumors and TP53 mutation frequencies across the breast tumor intrinsic subtypes for two different neoadjuvant treatment regimens: doxorubicin monotherapy (Doxo) and FUMI. Tumors are ordered according to the subtypes as presented in Fig. 1 in Sorlie et al. (8). Orange, tumor samples included in this study. TP53 mutation status is shown as percentage of tumors from both series combined (the normal breast-like subtype is excluded). Note that one progressive disease tumor from the FUMI series (Norway FU07-BE) and one stable disease tumor from the doxorubicin series (Norway 80-BE) were unclassified (uc). Source: PNAS, July 8, 2003, vol. 100, no. 14, 8418-8423. Copyright (2003) National Academy of Sciences, U.S.A.
treated with FUMI, 1 of 11 of the luminal A tumors versus 6 of 19 tumors from the other groups experienced progressive disease \( (P = 0.2146; \text{both data sets combined: } P = 0.0089) \). Interestingly, both of the luminal A type tumors expressing chemoresistance were wild-type for TP53. Although a luminal B profile was associated with resistance to doxorubicin (five of eight progressive disease; \( P = 0.0078) \), a similar finding was not identified among the tumors treated with 5-fluorouracil/mitomycin. A differential variation in response across subtypes has also been shown in a recently published study on breast cancer treated with preoperative chemotherapy (35, 37). Our finding that progressive disease was a rare event among tumors expressing the luminal A gene profile is interesting; however, these tumors rarely harbor mutations in the TP53 gene \( (6 \text{ of } 36 \text{ versus } 32 \text{ of } 44 \text{ among the other tumors; } P < 0.0001) \), so this may simply reflect TP53 status among these tumors. Nevertheless, this points to the importance of considering the molecular heterogeneity of tumors when assessing predictive as well as prognostic markers.

**Prediction of Therapeutic Response Classes Using Supervised Analyses**

To search the gene expression data for patterns associated with response (progressive disease, stable disease, or partial response; no complete response was recorded in these two series) and to explore the feasibility of using such patterns as predictors, PAM was used on all tumor samples obtained before therapy for each of three treatment groups: doxorubicin and FUMI separately and combined.

**Doxorubicin.** Training a predictor for progressive disease versus partial response (for which significant differences in gene expression might be expected) resulted in overall accuracy of 70% but with only three of eight progressive disease tumors correctly classified. When combining the response groups partial response and stable disease into one class, training of a predictor resulted in a similar accuracy (73%), now with five of eight progressive disease tumors correctly classified.

**5-Fluorouracil and Mitomycin.** Prediction of progressive disease versus partial response showed an accuracy of 78% with five of seven progressive disease tumors correctly classified. Next, training a predictor for progressive disease versus the combined groups partial response/stable disease resulted in an accuracy of 63% with only two of the progressive disease tumors correctly classified by cross-validation.

**Doxorubicin and FUMI Combined.** Similar PAM analyses for the two series combined (81 patients of whom 15 experienced progressive disease) showed similar accuracy rates; 62% for progressive disease versus partial response (6 of 15 progressive disease correctly classified) and 62% for progressive disease versus the partial response/stable disease combined group, with 7 progressive disease tumors correctly classified.

In addition to PAM, several additional statistical methods were used to determine if the less than optimal prediction accuracies were due to a particular analysis method (i.e., PAM). In particular, Random Forest, which is a multistep method for classification, and predictive modeling using a support vector machine method, termed r-SVM, which implements recursive gene ranking and selection steps, were both tested. All methods gave similar results, and thus, these results cannot be attributed to the statistical method used. A complete listing of the different methods and the prediction accuracies, sensitivities, and specificities resulting from the altogether seven prediction methods is presented in Supplementary Table S2.8 The values varied to some degree in magnitude, depending on the analysis method used and the sorting of the response groups. In particular, all predictors tended to do poorly in identifying the progressive disease tumors and often classified non-progressive disease samples correctly. This finding is a critical feature for the objective assessment of predictive profiles because, when a minor class is compared with a major class, a given “accurate” predictor could be developed that simply predicts most of the samples to be the major class.

The expression data used in this study were generated using several different production batches of cDNA arrays, and inconsistencies in such data that arose from process errors have been detected (38). Thus, we analyzed separately data from patients with progressive disease versus those with partial response using samples that had been hybridized on microarrays from the same batch only. Although this improved prediction accuracy up to 80%, only half of the progressive disease tumors were correctly classified (Supplementary Table S2).8 This finding of an inability to accurately identify most progressive disease tumors was true for all the predictors developed using the seven different methods, suggesting that this is inherent in the data and not due to the analysis method.

**Conclusions**

The aim of this study was to explore whether an analysis of gene expression data in a breast cancer cohort previously shown to yield prognostic gene profiles could identify gene signatures associated with response or resistance to chemotherapy. If so, this could add to the predictive value of TP53 mutations previously reported in the same tumors. However, we could not identify a gene profile using multiple diverse supervised analysis methods, which was highly accurate at identifying either drug-sensitive or drug-resistant tumors. Molecular tumor subtype was modestly correlated with response with luminal A tumors showing a lower rate of progressive disease and luminal B tumors treated with doxorubicin showing a high rate of progressive disease. In conclusion, we were not able to show that gene expression profiles can be used to accurately predict chemotherapy response in this data set. Similar to other studies (20–22, 24, 39), these results indicate that supervised analyses of relatively small sample sizes and with incomplete validation may not reveal a gene profile of sufficient predictive power to be of clinical use and suggest that genomic analyses using microarrays may need a different approach that incorporates functional hypotheses (40) to predict therapy sensitivity.
Prediction of Response in Breast Cancer

Acknowledgments

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References

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