Research article

Gene expression profiles derived from fine needle aspiration correlate with response to systemic chemotherapy in breast cancer

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Introduction

Chemotherapy given as adjuvant therapy after surgery to patients with primary operable breast cancer reduces the subsequent risk of relapse and death [1]. Although the absolute reduction in mortality is only about 10–12%, most patients now receive adjuvant chemotherapy because it is not possible to identify, at the start of treatment, which patients might gain benefit. Our approach to this problem

COX = cytochrome c oxidase; FNA = fine needle aspiration; HMG1 = high-motility group protein 1; PCR = polymerase chain reaction; RT = reverse transcriptase.
has been to give the chemotherapy before surgery (neoadjuvant therapy) to use the response in the primary tumor as a surrogate marker of subsequent survival benefit [2]. Predictions of response have been attempted on the basis of tumor expression of proliferation and apoptosis markers [3], endocrine and growth factors, and oncogenes [4]. However, no single tumor marker has been shown to possess a sufficient predictive value to render it clinically useful. To achieve a greater predictive value, multiple markers need to be examined and correlated with the response of tumor cells to chemotherapy. The development of cDNA microarray technology has provided such an opportunity. With this technology it has been possible to identify new classes in breast cancer according to their gene expression patterns and to correlate them with distinct clinical outcomes [5,6]. We have demonstrated previously that RNA amplification is suitable and reliable for the determination of gene expression profiles by using small amounts of RNA from core biopsies [7].

Our goal in this project was to determine the feasibility of obtaining representative cDNA expression array profiles from fine needle aspirations (FNAs) performed on breast carcinomas, which are known to have substantial heterogeneity after RNA amplification. We also evaluated the correlation between such profiles and subsequent clinical responses after neoadjuvant chemotherapy to determine whether this might provide a useful approach to response prediction for testing in future prospective studies.

Patients and methods

Patients
All patients provided informed consent before any procedures. To evaluate the sensitivity and reproducibility of the FNA technique, repeat presurgical FNA samples were taken from six patients. For two patients in this group, corresponding surgical specimens were available. Additionally, a group of 10 patients with breast cancer, who were treated with neoadjuvant chemotherapy, were evaluated by using three FNAs, two performed one week apart and before chemotherapy (adriamycin 60 mg/m² and cyclophosphamide 600 mg/m²), and the third on day 21 after the first cycle of chemotherapy. A 23-gauge needle (Oncotech Inc, Irvine, California, USA) was used for FNA in all cases.

RNA sample and cDNA preparation
From each aspirate a single-cell suspension was made in 2.5 ml of normal saline. Each sample was snap-frozen in liquid nitrogen and stored at –80°C. All FNA samples contained more than 50% tumor cells as assessed by a cytological examination of cytospin. The phenol/chloroform procedure (Trizol®; Gibco, Grand Island, New York, USA) was used to extract total RNA from each FNA sample. Total RNA isolated from the MCF10A human mammary epithelial immortalized cell line served as a common reference. Two rounds of Eberwine’s RNA amplification procedure were performed, with minor modifications, with total RNA from tumor FNA specimens and the MCF10A human mammary epithelial immortalized cell line as described previously [8]. Labeled cDNAs were prepared from 3 µg of amplified RNA, hybridized onto a 7600-feature glass cDNA microarray (NCI) and scanned with a 10 µm resolution GenePix 4000 scanner (Axon Instruments, Inc, Foster City, California, USA) as described elsewhere [9].

Data analysis
Expression profiles were analyzed with BRB Array tools, version 1.03 (Molecular Statistics and Bioinformatics Section, National Cancer Institute, Bethesda, Maryland, USA). Only spot-filtered data were considered for the analysis as described in the Supplementary Material. To compare gene expression profiles generated from each FNA–FNA and FNA–surgical specimen pair, we performed an unsupervised hierarchical cluster analysis [10]. A compound covariate predictor algorithm was applied to find a set of predictors for classifying each patient into the two predetermined classes (good responders versus bad responders) [11]. A cross-validated approach was performed to validate the class prediction: one patient was removed; the classifier was trained on the remaining patients and then tested for its ability to classify the withheld patient. To estimate the probability of obtaining the predicting membership purely by chance we performed random groupings of patients and we calculated the number of misclassifications (see Supplementary Material).

Real-time quantitative RT–PCR
Gene expression was measured with the GeneAmp 5700 Sequence Detector (Applied Biosystems, Foster City, California, USA). Primers and probes (BioServe Biotechnologies, Laurel, Maryland, USA) were designed with Primer Express software (see Supplementary Material) (Applied Biosystems). The following mRNAs were evaluated: CD44, HMG1, COX17, and ACTB as a normalizing control. Gene expression in each patient sample was then compared with expression in the reference cell line, MCF10A.

Results
FNAs can be used to obtain representative and reproducible gene expression profiles of breast tumors
To prove the feasibility of using RNA amplification we compared the gene expression profiles generated from each FNA with those obtained from the corresponding surgical specimen by using the unsupervised hierarchical clustering technique [10]. Two FNA samples out of 14 were rejected because of poor RNA quality (patients 11 and 13). All genes selected after spot filtering (4803 genes) were included in the clustering algorithm (See Supplementary Material). FNA–surgical specimen and FNA–FNA pairs from the same patient clustered together and were closer to each other than any samples from any other patients (Fig.1a). Similar results were obtained
when different distance-based clustering methods were used (data not shown).

Scatter plots were employed to investigate the level of similarity between the cDNA microarray results derived from various samples (Fig. 1b). When profiles from two FNAs from the same patient (obtained 1 week apart) were assessed (Fig. 1b, panels a and b), their Pearson correlation coefficient was very high ($r = 0.88$ and 0.94). This correlation is comparable to that observed between repeated hybridizations that we conducted with the same RNA ($r = 0.90–0.94$, data not shown). The comparison between surgical specimen and the corresponding FNAs also revealed a good correlation (Fig. 1b, panels c and d), albeit with lower correlation coefficients ($r = 0.74, 0.70$). In contrast, FNA–surgical specimen and FNA–FNA samples from different patients, as expected, gave distinctly different expression profiles with Pearson correlation coefficients of $r = 0.38$ and $r = 0.57$, respectively (Fig. 1b, panels e and f).

**Pretreatment gene expression profiles are associated with response or lack of response to chemotherapy in breast carcinoma**

To assess whether cDNA microarray profiles obtained from FNAs before chemotherapy would be able to differentiate and associate with the response or non-response to treatment, we studied an additional group of 10 patients with breast cancer who were to receive neoadjuvant chemotherapy. Patient demographics are shown in Table 1. Three pretreatment FNA samples out of 20 (patients 1, 3 and 4) were discarded because of an insufficient percentage of tumor cells. For each gene the average log expression ratio (ratio of the fluorescent intensities of the spots reflecting the relative level of gene expression) from the cDNA microarray experiments obtained from the two independent pretreatment FNAs was considered for analysis. Thirty-seven genes whose expression was most significantly different between good and poor responders were identified ($P < 0.01$; individual $t$-test). These included genes involved in the promotion of cell death, and chemosensitivity. To define a prediction model for classifying each patient into the two predetermined classes (good responders versus bad responders) we used a compound covariate predictor algorithm based on the expression levels of these 37 genes (Fig. 2a). Using this compound covariate predictor set of genes and a cross-validated approach we were able to correctly identify the response status of all 10 patients. Using 10,000 random permutations we confirmed that chance contribution to this prediction model was unlikely ($P < 0.009$) (see the Methods section and Supplementary Material).

**Early gene expression changes after one cycle of chemotherapy**

We also investigated the changes in gene expression after one cycle of chemotherapy. For this purpose we com-

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**Figure 1**

(a) Dendrogram representing similarities in the gene expression profiles between different FNAs and surgical specimens. FNA–surgical specimen and FNA–FNA pairs from the same patient clustered together more than with samples from any other patient. *Surgical specimen.

(b) Representative scatter plots indicating levels of similarity between the cDNA microarray results comparing different specimens. (panels a and b) Comparison of log expression ratios derived from two pretreatment FNAs from the same patient. (panels c and d) Comparison of log expression ratios derived from surgical specimens and the corresponding FNAs from two different patients. (panels e and f) Comparison of log expression ratios derived from two pretreatment FNAs and from a pretreatment FNA and a surgical specimen from different patients, respectively. The correlation coefficient of each comparison is shown at the top left of each panel.
pared the gene expression of the FNA taken on day 21 after the first cycle of chemotherapy with the average gene expression of the two pretreatment samples available for each patient. One post-treatment FNA out of 10

Table 1

| Patient no. | Age (years) | Tumor size (mm) | Node status | ER status | PgR status | Clinical response | Pathological finding (after chemotherapy) |
|-------------|-------------|-----------------|-------------|-----------|------------|------------------|-------------------------------------------|
| 1           | 42          | 7               | +           | +         | n/a        | CR               | No surgery                                |
| 2           | 37          | 50              | +           | +         | n/a        | MRD              | No surgery                                |
| 3           | 50          | 19              | +           | +         | n/a        | MRD              | DCIS only                                 |
| 4           | 56          | 24              | –           | –         | n/a        | MRD              | No surgery                                |
| 5           | 56          | 50              | +           | –         | n/a        | MRD              | DCIS only                                 |
| 6           | 48          | 50              | +           | –         | n/a        | SD               | RID                                       |
| 7           | 64          | 40              | +           | –         | –          | PR               | RID                                       |
| 8           | 47          | 30              | –           | +         | n/a        | SD               | RID                                       |
| 9           | 51          | 50              | –           | +         | n/a        | PR               | RID                                       |
| 10          | 67          | 30              | –           | –         | –          | SD               | RID                                       |

CR, complete clinical response (no residual palpable disease); ER, estrogen receptor; MRD, minimal residual disease (residual palpable irregularity that was too small to be measured); n/a, not available; PgR, progestrogen receptor; PR, partial response (more than 50% reduction in bidimensional measurement); SD, stable disease (between 50% reduction and 25% increase in bidimensional measurement); RID, residual invasive disease; DCIS, ductal carcinoma in situ.

Figure 2

Gene expression profiles distinguishing good from poor responders. (a) Hierarchical clustering of 37 genes that defined the class predictor and whose expression best differentiated good from poor responders before chemotherapy. Each row represents a single gene and each column represents the average of two available independent FNAs. Green and red squares indicate, respectively, overexpressed and underexpressed genes in a breast tumor compared with the MCF10A breast cancer cell line (color intensity is proportional to the magnitude of the expression level ratio). Black squares indicate genes with approximately equivalent expression levels and gray squares indicate missing or filter-excluded data. Branches representing good responders are shown in blue and those representing poor responders in yellow. (b) Hierarchical clustering of 16 genes whose change in expression best differentiated good from poor responders after one cycle of chemotherapy.
was discarded because of insufficient initial material (patient 1). In this analysis, outliers were defined as genes that showed at least a twofold change in their expression level (post-treatment compared with average pretreatment) in at least three out of five patients in each group. This was performed to select genes with consistently altered expression within each group. Using these stringent criteria, the numbers of outliers in the good responders were more than 10-fold those in the poor responder group (56 versus 5, respectively). When we used less stringent gene selection criteria based on the intersect of the genes that appeared at least once as an outlier, the good responders again exhibited much greater changes in gene expression than the poor responders (data not shown).

To further identify early gene expression changes that best discriminated the two groups after one cycle of chemotherapy, we also performed a parametric t-test on each gene’s log-ratio change value from each patient. As shown in Fig. 2b, we identified 16 genes whose change in expression level best differentiated between the two groups (P < 0.01, individual t-test).

**RT–PCR confirmation of microarray findings of selected genes**
To validate our array data, we performed RT–PCR of selected genes (COX17, HMG1 and CD44) involved in different molecular pathways identified in Fig. 2a, in the remaining RNA of all available pretreatment FNA samples. As shown in Fig. 3, the RT–PCR experiments confirmed the relative abundances of each of the three tested genes between good and poor responders obtained in microarray experiments.

**Discussion**
This study is the first to test the feasibility of obtaining representative array profiles from FNAs of breast carcinomas in a clinical setting. Amplification of tumor RNA from FNAs was used to achieve adequate signal. Our results demonstrate that when amplified RNA is used as a template, array profiles derived from an FNA of a given tumor are highly reproducible and representative. In addition, the molecular profiles generated from FNAs of different breast tumors can be distinguished from one another. Although the profiles derived from the surgical specimen and the corresponding FNAs showed discernible differences, these discrepancies are in the same range as the histologic differences between FNAs and surgical specimens [2]. Specifically, FNAs are enriched for the less adherent carcinoma cells and reduced in the stromal components in comparison with the intact tumor. Therefore, the profile of FNA samples might be more representative of the biology of the pure cancer cells because the stromal components have been removed. Most importantly, however, the composite expression profiles remain intact with repeat FNAs, such that comparisons between FNAs can be used to distinguish between different forms of breast tumor.

Our results also suggest that gene expression profiles obtained before and changes after one cycle of chemotherapy correlate with response to treatment. In the analysis of the pretreatment expression profiles, we identified 37 genes differentially expressed in pretreatment samples that together could segregate excellent responders from the poorer responders. Although the ‘leave one out’ cross-validation approach that we performed with the profiles from the pretreatment samples suggested that these 37 genes could correctly classify all patients, our numbers are small and the results should therefore be considered exploratory. Nevertheless, the comprehensive nature of array analysis permits the framing of hypothetical gene sets even with small sample sizes that can be vali-
dated with an independent and larger set of breast cancer patients in a prospective manner.

The change in gene expression as measured by the number of outliers when comparing the pretreatment and post-treatment samples seems to be more substantial in the responder than the non-responder group. This might be expected because changes in expression should be associated with the more effective cellular intervention. Thus, responders and non-responders could be differentiated both by specific gene changes and by the quantity of the change in expression.

An additional benefit of studying gene expression profiles is the identification of genes and/or gene pathways, which might associate with the response or intrinsic resistance to chemotherapy. Examination of the subset of genes identified before any treatment (Fig. 2a) revealed that these ‘discriminators’ are involved in a variety of cellular functions. This list included transcription factors (ERG and NFYB), oncogene (DLC1), growth factor receptors (KIT), genes involved in the ubiquitin–proteasome pathway (UBPH) and DNA repair and cell death regulators (HMG1, COX17, PAPPA, BCL2-like2, GADD34, RPL27, and CD44).

One of these genes, HMG1 (for high-motility group protein 1), was more highly expressed in responders. This gene encodes a structure-specific HMG-domain protein that is evolutionarily conserved. HMG1 binds preferentially to cruciform DNA, cisplatin-modified DNA, and other distorted structures. Overexpression of HMG1 has been shown to sensitize cells to cisplatin and carboplatin by shielding its major DNA adducts from nucleotide excision repair [12]. Depleting HMG1 and HMG2 from cell extracts by immunoprecipitation enhances the excision repair of cisplatin-modified DNA. Furthermore, introducing HMG2 by transfection enhances the cisplatin sensitivity of a lung adenocarcinoma cell line [13]. Thus, overexpression of HMG1 in the group of good responders might confer an increased sensitivity to the cytotoxic effect of chemotherapy.

Another interesting finding was the upregulated expression of COX17 (for cytochrome c oxidase assembly protein) in the good responders group. This gene encodes a copper-binding chaperone for cytochrome c oxidase (COX), the terminal enzyme of the mitochondrial respiratory chain. Dysfunction of the mitochondrial respiratory chain has been associated with apoptosis induction [14]. Interestingly, one of the cytotoxic mechanisms of doxorubicin is to inhibit COX activity [15]. Additionally, decreased COX expression was observed in doxorubicin-resistant leukemia K562 cells [16]. Thus, the overexpression of this cytochrome oxidase would be consistent with the association with better responders.

Finally, CD44, a main cell-surface receptor for hyaluronic acid, was downregulated in the group of good responders. CD44 is involved in matrix adhesion, lymphocyte activation, and the regulation of tumor proliferation. Several reports have shown an association between decreased CD44 expression and facilitation of apoptosis [17]. One of the mechanisms of this activity is thought to be the CD95/FAS-triggered induced shedding of CD44. Clinically, a decrease in CD44v6 expression has also been linked to a better response of neoadjuvant anthracycline-based treatment in cervical cancer [18].

Similar observations could also be made about genes whose expression was altered after one cycle of chemotherapy. For example, Cdk9, which encodes a catalytic subunit of TAK (cyclin T1/P-TEFb) and has been shown to have an antiapoptotic function during monocyte differentiation [19], was induced in the group of poor responders after the first treatment. TIMP-1, a collagenase inhibitor that was downregulated in the good responders, has been reported to control the cell growth phenotype during breast cancer development in an autocrine and paracrine manner [20]. Finally, DCTD, a gene that transcribes dCMP deaminase and is associated with increased resistance to cytarabine [21], was also induced in the poor responders. Taken together, these results indicate that the induction of genes associated with cellular resistance to chemotherapy seems to be associated with a poor clinical response, whereas a marker of cell growth seems to be downregulated in the good responders after one cycle of chemotherapy.

Our results indicate the complexity of genetic alterations involving various molecular pathways that might be associated with intrinsic and/or acquired resistance to chemotherapy in breast cancer. However, these expression profiles could represent a distinct signature for drug resistance in this particular small group of breast cancer patients. Interestingly, we have found that the dynamic changes after a single cycle of chemotherapy might further distinguish good responders from poor responders. The small numbers in our study preclude the definitive assignment of association genes with response, but these results have identified a significant list of plausible candidate markers that should be validated in larger clinical data sets.

In summary, our study demonstrates that expression profiles can be obtained in a reproducible manner from the small amount of RNA obtained from breast tumor FNAs. Moreover, we provide a proof of principle that profiles derived before treatment and changes in profiles shortly after starting treatment might have the potential to predict clinical outcomes from anthracycline-based chemotherapy in individual patients. Furthermore, using this approach we might be able to identify candidate genes and pathways,
which could be involved in intrinsic chemoresistance and could prove to be useful in identifying targets for the treatment of drug-resistant carcinomas.

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Supplementary material: patients and methods

Patients and collection of samples

At least three cytopsins were used for each FNA sample with the Shandon Cytopsin technique (Shandon, Pittsburgh, Pennsylvania, USA) on 0.5 ml of the cell suspension centrifuged at 500 rev/min for 5 min on 3-amino-propyltriethoxsilane slides. One of these slides was stained with May–Grünwald–Giemsa as a representative cytodiagnosis of the remaining 2 ml of the cell suspension to be used for RNA extraction. The remaining two slides were used to determine estrogen receptor and progesterone receptor status receptors by immunohistochemistry. RNA quality from each FNA was assessed by visualization of the ratio of 28S to 18S ribosomal RNA on a 1% agarose gel. Clinical response was defined as complete or nearly complete response (good response) compared with partial or no response (poor response), which we have previously defined as the best prediction model for subsequent relapse or survival advantage and used as the criteria for response in our previous marker studies [2].

Data analysis

Spot filtering

For each fluorescent channel, the log ratios were normalized by subtracting the median log ratio from each experiment. Spots of any experiment that had a red and/or green intensity less than 250 and 250 were flagged for absence of signal or no signal (poor response), which were flagged for any experimental reason in more than 10% of the microarray experiments were filtered out.

Compound covariate predictor

Classification of specimens into one of two predetermined classes based on gene expression data was performed with a compound covariate predictor. The predictor was built in two steps. First, a standard two-sample t-test was performed to identify genes with significant differences (at level a) in log expression ratios between the two classes. Second, the log expression ratios of differentially expressed genes were used as a basis for the single compound covariate [22] for each specimen; the compound covariate was used as the basis for defining a single compound covariate. The coexpression covariance for specimen i was defined as

$$C_i = \sum_{ij} X_{ij}$$
where $t_j$ is the $t$-statistic for the two-group comparison of classes with respect to gene $j$, $x_{ij}$ is the log ratio measured in specimen $i$ for gene $j$, and the sum is over all differentially expressed genes. Then the specimen to be classified was removed from the data set and the remaining specimens (comprising the training set) were used to determine the differentially expressed genes between the two classes. Using the log expression ratios of these genes, the value of the compound covariate was computed for every specimen in the training set and a classification threshold was calculated (the midpoint of the means of the compound covariates for the two classes was used as the threshold). The class of the omitted specimen was then predicted by computing the value of the compound covariate for the specimen, and which side of the threshold it fell on (that is, which class mean it was closest to) was determined. The entire process was repeated so that every specimen was omitted once and its class membership was predicted; the number of misclassified specimens was tallied. To determine whether the accuracy for predicting membership of specimens in the two given classes (as measured by the number of correct classifications) was better than the accuracy that could be attained for predicting membership in random groupings of specimens, we examined the distribution of the number of misclassifications for data sets in which the class labels were permuted. Random data sets were created by permuting class labels between the specimens, and cross-validated class prediction was performed on the resulting data sets as described above. The percentage of permutations that resulted in no greater number of misclassifications than for the original labeling of specimens was reported. If fewer than 5% of the permutations resulted in as few or fewer misclassifications, the accuracy of prediction into the given classes was considered significant.

**RT–PCR**

Primers and probes (BioServe Biotechnologies, Laurel, Maryland, USA) were designed with Primer Express software (Applied Biosystems). Probes were 5′-labeled with 6-carboxyfluorescein (FAM) and 3′-labeled with 6-carboxytetramethylrhodamine (TAMRA). Sequences were as follows. CD44: sense, 5′-TCACCAAATGCACCATTTCCT-3′; antisense, 5′-GTGATCAACAGTGCAATGGA-3′; probe, 5′-ACTTGCTGGCCTCTCGTGAGTCC-3′. COX17: sense, 5′-CGAGGCTGGCATAGATTTGG-3′; antisense, 5′-AACCAGACCCGGCATCTTTTG-3′; probe, 5′-TGTCTC-CGCTCATAGCTGCTTTTG-3′. HMG1: sense, 5′-GCGGACAAGGCCCCGTTA-3′; antisense, 5′-AGAGAAAGAA-GGCCGAAGGA-3′; probe, 5′-AAAAGATTCAAGGATCCCAATGCACCCC-3′. cDNA was generated from amplified RNA samples (2 ng per assay) by using MultiScribe reverse transcriptase and random hexamers for priming (Applied Biosystems). Reaction conditions were 10 min at 25°C, 30 min at 48°C, and 5 min at 95°C. PCR reactions were conducted in a volume of 25 µl by using AmpliTaq Gold DNA polymerase (Applied Biosystems), with sense and antisense primer concentrations of 900 nM each and a probe concentration of 250 nM. Thermal cycler parameters were 2 min at 50°C, 10 min at 95°C, and 40 cycles each consisting of a 15 s denaturing step at 95°C and a 1 min annealing/extension step at 60°C. All assays were performed in triplicate.