Breast Cancer Molecular Subtypes Respond Differently to Preoperative Chemotherapy

Roman Rouzier^{1,4} Charles M. Perou⁵ W. Fraser Symmans² Nuhad Ibrahim¹ Massimo Cristofanilli¹

View metadata, citation and similar papers at core.ac.uk

brought to you by CORE

Abstract Purpose: Molecular classification of breast cancer has been proposed based on gene expression profiles of human tumors. Luminal, basal-like, normal-like, and erbB2+ subgroups were identified and were shown to have different prognoses. The goal of this research was to determine if these different molecular subtypes of breast cancer also respond differently to preoperative chemotherapy.

Experimental Design: Fine needle aspirations of 82 breast cancers were obtained before starting preoperative paclitaxel followed by 5-fluorouracil, doxorubicin, and cyclophosphamide chemotherapy. Gene expression profiling was done with Affymetrix U133A microarrays and the previously reported "breast intrinsic" gene set was used for hierarchical clustering and multidimensional scaling to assign molecular class.

Results: The basal-like and erbB2+ subgroups were associated with the highest rates of pathologic complete response (CR), 45% [95% confidence interval (95% Cl), 24-68] and 45% (95% Cl, 23-68), respectively, whereas the luminal tumors had a pathologic CR rate of 6% (95% Cl, 1-21). No pathologic CR was observed among the normal-like cancers (95% Cl, 0-31). Molecular class was not independent of conventional clinicocpathologic predictors of response such as estrogen receptor status and nuclear grade. None of the 61 genes associated with pathologic CR in the basal-like group were associated with pathologic CR in the erbB2+ group, suggesting that the molecular mechanisms of chemotherapy sensitivity may vary between these two estrogen receptor – negative subtypes.

Conclusions: The basal-like and erbB2+ subtypes of breast cancer are more sensitive to paclitaxel- and doxorubicin-containing preoperative chemotherapy than the luminal and normal-like cancers.

© 2005 American Association for Cancer Research. doi:10.1158/1078-0432.CCR-04-2421

Breast cancer is a clinically heterogeneous disease. Histologically similar tumors may have different prognoses and may respond to therapy differently. It is believed that these differences in clinical behavior are due to molecular differences between histologically similar tumors. DNA microarray technology is ideally suited to reveal such molecular differences. A novel molecular classification of breast cancer based on gene expression profiles was recently proposed (1). The investigators identified a set of stably expressed genes ("intrinsic gene set"; n = 534) that accounted for much of the molecular differences between 42 breast cancers and did hierarchical cluster analysis to identify subgroups of cancers with separate gene expression profiles. Luminal, basal-like, normal-like, and erbB2+ subgroups were identified and were shown to have different prognoses (1-4). These results were confirmed in follow-up experiments by the same group and others using larger numbers of cases. The basal-like (mostly estrogen receptor negative) and erbB2+ (mostly HER-2 amplified and estrogen receptor negative) subgroups had the shortest relapse-free and overall survival, whereas the luminal-type (estrogen receptorpositive) tumors had a more favorable clinical outcome (2-4). There is no published data on how the different molecular

Authors' Affiliations: Departments of ¹Breast Medical Oncology, ²Pathology, and ³Biostatistics and Applied Mathematics, University of Texas M.D. Anderson Cancer Center, Houston, Texas; ⁴Unité Propre de l'Enseignement Supérieur EA 3535, Institut Gustave Roussy, Villejuif, France; ⁵Lineberger Comprehensive Cancer Center, University of North Carolina, Chapel Hill, Noth Carolina; ⁶Millennium Pharmaceuticals, Inc., Cambridge, Massachusetts; and ⁷Albany Medical College, Albany, New York

Received 11/24/04; revised 4/4/05; accepted 5/18/05.

Grant support: The Nellie B. Connally Breast Cancer Research Fund, Millennium Pharmaceuticals, The Dee Simmons Fund, University of Texas M.D. Anderson Cancer Center Aventis Drug Development Award (L. Pusztai), The Susan G. Komen Breast Cancer Foundation (grant LF2002-044HM; W.F. Symmans), Association pour la Recherche sur le Cancer (R. Rouzier), and National Cancer Institute Specialized Program of Research Excellence in Breast Cancer (grant P50-CA58223-09A1; C.M. Perou).

The costs of publication of this article were defrayed in part by the payment of page charges. This article must therefore be hereby marked *advertisement* in accordance with 18 U.S.C. Section 1734 solely to indicate this fact.

Note: Supplementary data for this article are available at Clinical Cancer Research Online (http://clincancerres.aacrjournals.org/).

Requests for reprints: Lajos Pusztai, Department of Breast Medical Oncology, University of Texas M.D. Anderson Cancer Center, Unit 424, 1515 Holcombe Boulevard, Houston, TX 77030-4009. Phone: 713-792-2817; Fax: 713-794-4385; E-mail: Jousztai@mdanderson.org.

classes of breast cancer respond to chemotherapy. The goal of the current project was to examine if these different molecular subclasses of breast cancer also respond differently to anthracycline- and paclitaxel-containing preoperative chemotherapy.

Patients and Methods

Fine needle aspirations of breast cancer were collected in a prospectively designed pharmacogenomic marker discovery study at the Nellie B. Connally Breast Center of the University of Texas M.D. Anderson Cancer Center. The goal of the ongoing clinical study was to develop multigene predictors of pathologic complete response (CR) to preoperative therapy. The current analysis was undertaken to examine if molecular class is associated with sensitivity to chemotherapy. Gene expression results from the first 82 patients with stage I to III breast cancer were included in this analysis. Patient characteristics were presented in Table 1. Fine needle aspiration was done using a 23- or 25-gauge needle before starting preoperative chemotherapy with 12 weeks of paclitaxel followed by 5-fluorouracil, doxorubicin, and cyclophosphamide \times 4 courses. Cells from 2 to 3 passes were collected into vials containing 1 mL of RNAlater solution (Ambion, Austin, TX) and stored at -80°C. Median RNA yield of the 82 specimens was 2.0 µg (1-22 µg). Approximately 70% of all aspirations vielded at least 1 µg total RNA, which is required for gene expression profiling. The main reason for failure to obtain sufficient RNA was acellular aspirations (low cell yield). The cellular composition of the fine needle aspiration samples was previously reported; in brief, fine needle aspiration samples on average contain 80% neoplastic cells and the rest of the cells are infiltrating leukocytes (5). These samples contain little or no stromal cells (fibroblasts and adipocytes) or normal breast epithelium. Of the 82 RNA specimens used in this analysis, 33 were included in a previous pharmacogenomic analysis using cDNA arrays (6). These 33 cases were profiled on both platforms (Affymetrix U133A and proprietary cDNA) and the results of the cross platform comparison of gene expression data were published separately (7). All patients underwent surgery after completion of 24 weeks of preoperative chemotherapy. Grossly visible residual cancer was measured and representative sections were submitted for routine histopathologic examination. When there was no grossly visible residual cancer, the slices of the specimen were radiographed and all areas of radiologically and/or architecturally abnormal tissue were entirely submitted for histopathologic study. Patients without any residual invasive cancer in the breast and axillary lymph nodes were considered to have pathologic CR. Patients with residual in situ cancer (DCIS) only were also considered to have pathologic CR. Estrogen receptor and HER-2 status was determined by routine clinical diagnostic methods [using mouse monoclonal anti-estrogen receptor antibody 6F11 (Novacastra/Vector Laboratories, Burlingame, CA) and fluorescence in situ hybridization assay to determine HER-2 amplification (PathVision kit, Vysis, Dovners Grove, IL)] on a diagnostic core needle biopsy obtained before or concomitant to the research fine needle aspiration. Nuclear grade was defined by the modified Black's nuclear grading system (1 = low grade, 2 = intermediate grade, and 3 = high grade; ref. 8). The study was approved by the Institutional Review Board of M.D. Anderson Cancer Center, and all patients signed an informed consent.

RNA was extracted from fine needle aspiration samples using the RNAeasy Kit (Qiagen, Valencia CA). The amount and quality of RNA were assessed with a DU-640 UV Spectrophotometer (Beckman Coulter, Fullerton, CA) and by an Agilent 2100 Bioanalyzer RNA 6000 LabChip kit (Agilent Technologies, Palo Alto, CA). Profiling was done without second round amplification using a minimum of 1 μ g total RNA. Double-stranded cDNA was synthesized, followed by *in vivo* transcription reaction to generate biotinylated cRNA. Biotin-labeled and fragmented cRNA was hybridized to Affymetrix U133A gene chips

overnight at 42°C. The Affymetrix GeneChip system was used for hybridization and scanning and the dCHIP V1.3 (http://dchip.com) software was used to generate probe level signal and for normalization of data across arrays.

Data Analysis

dCHIP V1.3 software was used for normalization; this program normalizes all arrays to one standard array that represents a chip with median overall intensity. After normalization, estimates of feature level intensity were derived from the 75th percentile of pixel level intensity of each feature. Each individual probe was aggregated at the feature level to form a single measure of intensity for each probe set. We used the perfect match model. Statistical analysis was

 Table 1
 Clinical information and demographics of the

patients included in the study ($n = 82$)
Female	82 (100%)
Median age	52 y (range 29-79)
Race	
Caucasian	56 (68%)
African American	11 (13%)
Asian	7 (9%)
Hispanic	6 (7%)
Mixed	2 (2%)
Histology	
Invasive ductal	73 (89%)
Mixed ductal/lobular	6 (7%)
Invasive lobular	1 (1%)
Invasive mucinous	2 (2%)
Tumor-node-metastasis stage	
T ₁	7 (9%)
T ₂	46 (56%)
T ₃	15 (18%)
T ₄	14 (17%)
No	28 (34%)
N ₁	38 (46%)
N ₂	8 (10%)
N ₃	8 (10%)
Nuclear grade (benign melanocytic nevus)	
1	2 (2%)
2	23 (37%)
3	35 (61%)
Estrogen receptor positive*	35 (43%)
Estrogen receptor negative	47 (57%)
HER-2 positive [↑]	57 (70%)
HER-2 negative	25 (30%)
Neoadjuvant therapy ⁺	
Weekly T (80 mg/m ²) \times 12 + FAC \times 4	69 (84%)
3-weekly T (225 mg/m Cl) $ imes$ 4 + FAC $ imes$ 4	13 (16%)
Pathologic CR	21 (26%)
Residuald isease	61 (74%)

*Cases where >10% of tumor cells stained positive for estrogen receptor with immunohistochemistry were considered positive.

[†]Cases that showed gene copy number > 2.0 were considered HER-2 positive. [‡]T, paclitaxel; Cl, 24-hour continuous infusion; and FAC, 5-flurouracil (500 mg/m²), doxorubicin (50 mg/m²), and cyclophosphamide (500 mg/m²). done by using the BRB-Arraytools version 3.0 software package (http://linus.nci.nih.gov/BRB-ArrayTools.html). Complete linkage hierarchical clustering was done with the previously published breast cancer intrinsic gene set with 1-Pearson correlation coefficient as distance metric (1). Cluster reproducibility and the robustness of the dendograms were examined by the method proposed by McShane et al. (9) based on 500 perturbations. Tumors clustering together in significant dendrogram branches were categorized as one molecular class. We also used multidimensional scaling with the Eucledian distance as metric to provide graphical representation of the distances among samples. This method also made it possible to test global statistical significance to determine whether the expression profiles form distinct clusters (rather than represent the same multivariate Gaussian distribution). Genes differentially expressed in a particular molecular class compared with all other tumors and between cases of pathologic CR and residual cancer within a single molecular subgroup were identified using the significance analysis of microarrays (SAM) software with 1,000 sample permutations. SAM uses permutations to estimate the false discovery rate and an adjustable threshold allows for control of the false discovery rate (10).

Pathologic complete response rates were calculated for each molecular class and assessed in univariate analysis (χ^2 test) and multivariate analysis (logistic regression). Estrogen receptor and HER-2 status, nuclear grade, tumor size, and lymph node involvement were included in the multivariate analysis. We built logistic regression – based prediction models with various combinations of clinical variables and molecular class to examine if knowledge of the molecular class improves prediction accuracy above what can be achieved by combining routine clinical variables.

Results

Hierarchical clustering with the breast cancer intrinsic gene set reveals previously described molecular classes in fine needle aspiration specimens. The intrinsic breast cancer gene set consists of 534 genes of which expression showed significantly larger variation between tumors than between paired samples from the same tumor in an early seminal publication (1). Of these intrinsic genes, 424 were represented on the Affymetrix U133A chip. We did supervised hierarchical clustering with 689 Affymetrix probe sets that represented these 424 genes to define the molecular classes of breast tumors in our data. The tumors clustered into four major classes. The reproducibility indices of the four distinct clusters were 0.82, 0.76, 0.85, and 0.78, respectively, which indicates reasonably robust clusters (9). Tumors within each molecular subtypes corresponded well to the previously described clinicopathologic phenotypes of luminal (n = 30), normal-like (n = 10), basal-like (n = 22), and erbB2+ (n = 20) cancers (Fig. 1A). All of the luminal tumors were estrogen receptor positive by immunohistochemistry. All but two cases (80%) of the erbB2+ molecular class had HER-2 gene amplification by fluorescence in situ hybridization analysis. All but one of the basal-like tumors (95%) was estrogen receptor negative and 75% of these tumors were also high nuclear grade. These groups did not differ significantly in nodal status, tumor size, or patient age distribution (Fig. 1B). Multidimensional scaling analysis also confirmed the presence

of significant clustering of the cases (global test of significance P = 0.04, Fig. 1C). To examine how sensitive the cluster results are to the actual gene set used for clustering, we did a multidimensional scaling analysis using the probe sets with the highest variance (top 10%) across all samples (2,228 probe sets including 229 overlapping probes with the intrinsic gene set). Cases with the same molecular class (as defined by the intrinsic gene set) continued to cluster together (global test of significance P = 0.047; Fig 1E). This suggests that the gene signature – based groups are robust.

To define the molecular differences further between the subgroups, we identified differentially expressed genes between the four molecular classes using SAM analysis on the most variably expressed probe sets (n = 2,228). Setting the most stringent false discovery rate at 0.0001, 372 probe sets representing 298 genes were identified as differentially expressed between the four distinct groups (Supplementary Table S1). The high expression of estrogen receptor 1 and several of the known estrogen receptor-inducible genes, such as X-box binding protein 1 and SLC39A6 among many others, characterized the luminal subgroup. The basal-like subgroup was characterized by the expression of keratin 17, keratin 5, and γ -aminobutyric acid receptor π subunit among others. The erbB2+ subtype was characterized by the overexpression of genes that are located in the HER-2 amplicon including erbB2 and GRB7. Interestingly, the normal-like group had only 15 genes that were overexpressed in this subgroup. These gene lists could be used to further characterize the various molecular subclasses and for the development of supervised molecular class prediction methods.

Correlation between molecular class and pathologic complete response to preoperative chemotherapy. The rates of pathologic CR differed significantly among the four molecular classes of breast cancer defined by clustering using the intrinsic gene set. Basal-like and erbB2+ subgroups were associated with the highest rate of pathologic CR, 45% [95% confidence interval (95% CI), 24-68] and 45% (95% CI, 23-68), respectively, whereas luminal tumors had a pathologic CR rate of 6% (95% CI, 1-21). No pathologic complete response was observed in the normal-like subclass (Table 2). We next used multidimensional scaling graph to explore if the breast intrinsic gene set can separate cases with pathologic CR versus those with residual disease (Fig. 1D). The global test of significance showed that the observed clusters of pathologic CR and residual disease were significantly separate (P = 0.026).

Next, we examined the clinical pathologic variables that were associated with pathologic CR in this data. Age < 50 years and estrogen receptor-negative status were identified as independent variables associated with higher likelihood of pathologic CR in multivariate analysis including age, estrogen receptor and HER-2 status, tumor size, clinical nodal status, and nuclear grade (Table 3). To examine if knowledge of the molecular class improves estimation of probability of pathologic CR beyond what can be achieved with routine clinical variables, we built three different logistic regression models including the clinical variables (age, tumor, node stage), the histopathologic variables (grade, estrogen receptor, and HER-2 status), and the molecular class in various combinations. For this analysis, we merged the luminal and the normal-like groups because there was no pathologic CR in the normal-like category and these tumors were phenotypically similar to the luminal tumors



В

		Molecu	ilar class		
	Luminal	normal-like	erbb2	basal-like	
	n=30	n=10	n=20	n=22	p value
Age < 50	7 (23%)	4 (40%)	11 (50%)	7 (35%)	.72
T > 5cm	9 (30%)	5 (50%)	8 (36%)	7 (35%)	.68
N1-3	17 (57%)	7 (70%)	13 (59%)	15 (75%)	.57
ER+	30 (100%)	6 (60%)	11 (55%)	1 (5%)	<.001
HER+	1 (3%)	1 (10%)	16 (80%)	3 (14%)	<.001
High BNG	11 (37%)	3 (30%)	16 (80%)	20 (91%)	<.001



Fig. 1. Different molecular classes of breast cancer show distinct clinicopathologic features and respond differently to chemotherapy. *A*, hierarchical clustering using the 689 probe sets (424 unique genes) corresponding to the breast intrinsic gene set (n = 82 cases). *B*, correlation between molecular subclass and clinicopathologic characteristics in univariate analysis. *C*, graphical representation of the distances among the samples that belong to different molecular classes using multidimensional scaling (*P* = global test of significance for molecular subclasses). *D*, cases with pathologic CR form a separate cluster in multidimensional scaling analysis. *E*, when genes with the greatest variance (top 10%, n = 2228) across samples were used for multidimensional scaling, rather than the intrinsic gene set, molecular subgroups continued to cluster together.

Table 2.	Correlation b	between	molecular	classif	fication	and	patholo	gic o	complete i	response
----------	---------------	---------	-----------	---------	----------	-----	---------	-------	------------	----------

	Pathologic com	plete response	
	No	Yes	
Molecular classification	n [% (95% Cl)]	n [% (95% Cl)]	
Luminal A/B subtype	28 [93% (78-99)]	2 [7% (1-22)]	
Normal breast like	10 [100% (29-100)]	0 [0% (0-31)]	
erbB2+	11 [55% (32-77)]	9 [45% (23-68)]	
Basal subtype	12 [55% (32-76)]	10 [45% (24-68)]	<i>P</i> < 0.001

(HER-2 normal and estrogen receptor positive) that also had low pathologic CR rates. Molecular class was not independently associated with pathologic CR because of the high correlation between molecular class and estrogen receptor status and nuclear grade in this cohort. We constructed Receiver Operating Characteristic curves to measure the predictive accuracy of the logistic regression models including (*a*) clinical + pathologic variables, (*b*) clinical variables + molecular classification, and (*c*) clinical + pathologic variables + molecular class (Fig. 2). The three models yielded similar areas under the Receiver Operating Characteristic curve. This indicates that the molecular class alone can replace histopathologic characteristics (estrogen receptor, HER-2 status, or grade) for prediction of pathologic CR but provides little additional information when these characteristics are included. More directed supervised class prediction methods may be needed to develop a multigene predictor of pathologic CR. Such predictors can be developed by identifying informative genes that are differentially expressed between cases of pathologic CR and residual disease and combing these genes into a weighted prediction score or algorithm.

Table 3. Multivaria	te analysis of pro	edictive fa	ctors for pathologic	CR		
Variables	Model 1: clinical and histologic variables		Model 2: clinical variables and molecular classification		Model 3: clinical, histologic variables, molecular classification	
	OR (95% CI)	Ρ	OR (95% CI)	Ρ	OR (95% CI)	Р
Age (y)						
<50	1		1		1	
>50	0.27 (0.8-0.91)	0.035	0.17 (0.06-0.45)	< 0.001	0.43 (0.11-1.7)	0.43
Tumor (cm)						
<5	1		1		1	
>5	0.55 (0.14-2.3)	0.41	0.53 (0.15-1.8)	0.28	0.64 (0.14-2.9)	0.56
Node						
No	1		1		1	
N ₁₋₃	0.96 (0.31-3.0)	0.94	0.65 (0.22-2.0)	0.18	0.90 (0.22-3.7)	0.90
Estrogen receptor						
Negative	1				1	
Positive	0.12 (0.02-0.31)	<0.001			0.08 (0.02-0.35)	0.001
HER-2						
Negative	1				1	
Positive	1.77 (0.42-7.5)	0.43			0.32 (0.03-3.6)	0.34
Nuclear grade						
1/2	1				1	
3	2.6 (0.81-8.4)	0.11			2.5 (0.4-13.6)	0.30
Histology						
Ductal	1				1	
Other	1.14 (0.17-7.5)	0.89			2.3 (0.11-1.7)	0.76
Molecular classification						
Luminal/normal-like			1		1	
Normal-like			0 (0)	0.99	0 (0)	0.99
Basal-like			3.3 (1.0-11)	0.06	0.8 (0.12-5.5)	0.83
erbB2+			4.4 (1.2-17)	0.026	7.8 (0.62-100)	0.11

NOTE: Multivariate analysis of different combinations of clinical (age, tumor, node) and histopathologic characteristics (grade, estrogen receptor and HER-2 status, and histologic type) and molecular class as variables. Three distinct prediction models were examined: clinical plus histopathologic variables (model 1), clinical variables plus molecular class (model 2), and all three types of information together (model 3).

Genes associated with pathologic complete response in the different molecular subgroups. We next examined if the genes of which expression was associated with pathologic CR differed between basal-like and erbB2+ subtypes. Because only two cases of pathologic CR were observed in the luminal group and no pathologic CR was seen in the normal-like group, these groups were not included in this exploratory analysis. Seventytwo probe sets (corresponding to 61 genes) were differentially expressed between basal-like tumors that achieved a pathologic CR and those that did not (Table 4). All highly variable genes (n = 2,228) were used in this analysis and the false discovery rate was set at 5%. Interestingly, within the erbB2+ group, zero genes were identified at false discovery rate < 10%. If the false discovery rate was set at 50%, 16 probe sets (15 genes) were identified; however, half of these could represent spurious discovery (data not shown). A greater variance of gene expression among erbB2+ tumors and, therefore, greater molecular heterogeneity compared with basal-like tumors combined with the small sample size (erbB2+; n = 20) may explain why it was difficult to identify genes in this group. Importantly, none of the genes associated with pathologic CR in the basal-like group was associated with pathologic CR in the erbB2+ group. We also assessed if there was a correlation between fold differences of expression of pathologic CR-associated genes in cases with pathologic CR compared with residual disease in basal-like and erbB2+ tumors, respectively. There was no correlation (P =0.19). The absence of correlation suggests that genes associated with chemotherapy sensitivity are different between these two molecular subgroups of breast cancer.

Discussion

The goal of this current project was to further evaluate the clinical relevance of a novel gene expression–based classification system of breast cancer. This new classification is based on gene expression signatures of variably expressed genes in breast cancer (1). It has previously been shown that the various molecular classes have different long-term survivals (1-4). However, it is not possible to decipher from these earlier studies if the differences in survival are due to different metastatic potentials or to different sensitivities to adjuvant chemotherapy or hormonal therapy because the patients included in these studies received various forms of multi-

modality treatment. In the current study, we examined newly diagnosed stage I to III breast cancers that all received preoperative treatment with anthracycline and taxane followed by surgery to determine if the different molecular classes show different chemotherapy sensitivities based on pathologic response to preoperative chemotherapy.

All previous reports on molecular classification used frozen breast cancer tissues for gene expression profiling. The current study differs from these in that we used fine needle aspiration specimens. Surgically resected cancer tissues differ from fine needle aspiration in cellular composition. The fine needle aspiration material contains 80% to 90% pure neoplastic cells whereas surgical biopsies or core needle biopsies contain a variable amount of stromal cells. It was therefore of interest to determine if the intrinsic gene set that discriminated molecular class in surgical specimens could also separate molecular classes of breast cancer in fine needle aspiration data. If such separation can be observed, this would suggest that these informative genes are primarily expressed in neoplastic cells rather than in stromal cells.

In the current study, we did hierarchical clustering and multidimensional scaling analysis using the breast cancer intrinsic gene set which mimics the original class discovery process because there are presently no uniformly accepted class prediction tools to define the molecular classes of breast cancer utilizing gene expression data. We observed very similar results in our fine needle aspiration data as was reported by others on surgical tissues. The two most readily distinguishable molecular classes of breast cancer are the basal-like and luminal subtypes whereas the normal-breast like class is the least robust. This may be due to the possibility that the original samples in this category contained significant amount of contaminating normal breast tissue. The basal-like, erbB2+, and luminal subclasses were distinguished by some of the same genes and histologic phenotypes in our series as previously reported. This supports the hypothesis that these clusters represent genuinely different diseases within breast cancer (3, 4).

The different molecular classes of breast cancer showed different sensitivities to preoperative chemotherapy. The basallike and erbB2+ subgroups had the highest rates of pathologic CR, 45% (95% CI, 23-68). The luminal and normal-like tumors had low pathologic CR rates of 6% (95% CI, 1-21)





Table 4.	Genes	associated	with	pathologic	CR	in
basal-like	breast o	cancer				

	Probe set	Gene symbol	Fold difference of means between residual disease and pathologic CR
1	213060_s_at*	CHI3L2	4.117
2	200052_s_at	ILF2	2.645
3	213338_at	RIS1	3.181
4	214433_s_at*	SELENBP1	2.735
5	204319_s_at	RGS10	2.183
6	213005_s_at	ANKRD15	2.517
7	221561_at	SOAT1	2.156
8	212190_at	SERPINE2	2.448
9	209387_s_at	TM4SF1	3.096
10	221727_at	PC4	1.842
11	213716_s_at	SECTM1	2.183
12	203165_s_at	SLC33A1	1.826
13	207414_s_at	PACE4	1.732
14	201819_at	SCARB1	2.118
15	217983_s_at	RNASET2	1.915
16	214540_at	HIST1H2BO	1.864
17	218538_s_at	MRS2L	1.794
18	202506_at	SSFA2	1.678
19	215071_s_at	HIST1H2AC	2.099
20	202988_s_at	RGS1	2.328
21	220624_s_at	ELF5	2.521
22	221505_at*	ANP32E	2.51
23	208370_s_at	DSCR1	2.332
24	204407_at	TTF2	1.913
25	218398_at	MRPS30	1.595
26	213/54_s_at	IRIM26	1.844
27	210147_at	ARI3	3.943
28	204809_at	CLPX	1.813
29	202035_s_at	SFKFI	0.401
30 21	203303_x_al	CKS1R	1.752
37	201037_5_dl	URF2G1	1.337
52	200142_3_dl	ODLZUI	1.740

and 0% (95% CI, 0-21%), respectively. However, the pathologic predictors of response (i.e., grade and estrogen receptor status). The basal-like and erbB2+ tumors were predominantly high nuclear grade and the basal-like tumors were almost all estrogen receptor negative. Both of these characteristics are known to be associated with higher likelihood of pathologic CR to preoperative chemotherapy (11–13). Because of this association, incorporation of molecular class into a logistic regression–based predictor of response did not improve the prediction accuracy compared with using routine clinical and pathologic variables only. Therefore, it is likely that more focused gene signature–based predictors will need to be developed through supervised outcome prediction methods.

How to define the best multigene predictor of response to chemotherapy is not known. One approach is to group all breast cancers into either responders (e.g., pathologic CR) or nonresponders (e.g., residual disease), define the gene expression differences between these groups, and use this **Table 4.** Genes associated with pathologic CR inbasal-like breast cancer (Cont'd)

	Probe set	Gene symbol	Fold difference of means between residual disease and pathologic CR		
33	209340_at	UAP1	1.724		
34	203362_s_at	MAD2L1	1.967		
35	217028_at	CXCR4	2.097		
36	205044_at*	GABRP	6.224		
37	36711_at	MAFF	2.271		
38	202023_at	EFNA1	1.721		
39	212915_at	PDZRN3	2.539		
40	217851_s_at	C20orf45	1.734		
41	211762_s_at	KPNA2	1.849		
42	213134_x_at*	BTG3	2.007		
43	204162_at	KNTC2	2.283		
44	212276_at	LPIN1	2.219		
45	219768_at	B7-H4	1.843		
46	209551_at	MGC11061	1.925		
47	203744_at	HMGB3	1.457		
48	200975_at	PPT1	1.628		
49	221931_s_at	SEC13L	1.796		
50	209786_at	HMGN4	1.63		
51	218963_s_at	KRT23	3.088		
52	219209_at	MDA5	2.337		
53	214214_s_at	C1QBP	1.799		
54	209656_s_at	TM4SF10	2.645		
55	203706_s_at*	FZD7	2.603		
56	206055_s_at	SNRPA1	1.799		
57	204825_at	MELK	1.735		
58	212762_s_at	TCF7L2	1.928		
59	203423_at	RBP1	1.82		
60	210605_s_at*	MFGE8	2.085		
61	214835_s_at	SUCLG2	1.576		
NOT gene	NOTE: There are 72 probes sets significant by SAM, corresponding to 61 genes. The median false discovery rate among the 72 significant genes is 0.04. Cores or worked by significant				

machine learning-based predictor. This approach was successfully applied to develop prognostic signatures for breast cancer and was also promising in small pilot studies of chemotherapy response prediction (6, 14, 15). To develop the best possible supervised classifier for prediction of pathologic CR from this data set was not the goal of this current analysis. However, if distinct molecular classes of breast cancer exist, one could hypothesize that stratification of patients by molecular class may yield more accurate class-specific predictors than unstratified use of the data.

As an exploratory analysis, we attempted to define the molecular differences between tumors that are extremely chemotherapy sensitive (pathologic CR) and those that are more resistant (residual disease) within the basal-like and erbB2+ groups, separately. In the basal-like group (n = 22, including 10 pathologic CR), 61 genes were identified that were statistically significantly associated with pathologic CR. It is important to realize that none of these genes are associated with estrogen receptor status or high grade (the two

conventional strong predictors of pathologic CR) because the basal-like group almost exclusively consists of high-grade and estrogen receptor-negative tumors. We could not define a robust gene set that correlated with pathologic CR in the erbB2+ group (n = 20, including 9 pathologic CR). Importantly, the genes that were associated with pathologic CR in the basal-like group were not associated with pathologic CR in the erbB2+ group. This suggests that distinct sets of genes are associated with pathologic CR in the different molecular classes.

It is tempting to speculate on the biological function of the genes that are differentially expressed between cases with pathologic CR and those with residual cancer. However, not all of these genes may play a causative role in determining sensitivity to chemotherapy. Some of these may be distant downstream transcriptional effects of biological events that influence drug sensitivity and a few could represent spurious discovery. From the vantage point of gaining mechanistic insight into the biology of chemotherapy sensitivity or resistance, these gene lists should be regarded as hypothesisgenerating and will require further *in vitro* experimentation to show a functional role for any particular molecule.

In summary, these results indicate that the major molecular classes of breast cancer can be detected in gene expression data regardless of tissue sampling method (i.e., fine needle aspirations, core needle, or surgical biopsies). The different molecular classes of breast cancer not only have different prognoses but also show distinct sensitivities to preoperative chemotherapy. The basal-like and erbB2+ subtypes of breast cancer are more sensitive to paclitaxel- and doxorubicincontaining preoperative chemotherapy than the luminal and normal-like cancers. The genes associated with pathologic CR were different between the basal-like and erbB2+ subgroups, which suggest that the mechanisms of chemotherapy sensitivity may vary across the subtypes. The possibility that distinct predictive signatures can be developed for the different molecular subtypes of breast cancer warrants further examination.

References

- 1. Perou CM, Sorlie T, Eisen MB, et al. Molecular portraits of human breast tumours. Nature 2000;406: 747–52.
- Sorlie T, Perou CM, Tibshirani R, et al. Gene expression patterns of breast carcinomas distinguish tumor subclasses with clinical implications. Proc Natl Acad Sci U S A 2001;98:10869–74.
- Sorlie T, Tibshirani R, Parker J, et al. Repeated observation of breast tumor subtypes in independent gene expression data sets. Proc Natl Acad Sci U S A 2003;100:8418–23.
- Sotiriou C, Neo SY, McShane LM, et al. Breast cancer classification and prognosis based on gene expression profiles from a population-based study. Proc Natl Acad Sci U S A 2003;100: 10393–8.
- Symmans WF, Ayers M, Clark EA, et al. Total RNA yield and microarray gene expression profiles from fine-needle aspiration biopsy and core-needle biopsy samples of breast carcinoma. Cancer 2003;97: 2960–71.
- 6. Ayers M, Symmans WF, Stec J, et al. Gene expression

profiles predict complete pathologic response to neoadjuvant paclitaxel and fluorouracil, doxorubicin, and cyclophosphamide chemotherapy in breast cancer. J Clin Oncol 2004;22:2284–93.

- Stec J, Wang J, Coombes K, et al. Comparison of the predictive accuracy of DNA array based multigene classifiers across cDNA arrays and Affymetrix Gene-Chips. J Mol Diagnostics. In press 2005.
- Cutler SJ, Black MM, Mork T, et al. Further observations on prognostic factors in cancer of the female breast. Cancer 1969;24:653–67.
- McShane LM, Radmacher MD, Freidlin B, et al. Methods for assessing reproducibility of clustering patterns observed in analyses of microarray data. Bioinformatics 2002;18:1462–9.
- **10.** Tusher VG, Tibshirani R, Chu G. Significance analysis of microarrays applied to the ionizing radiation response. Proc Natl Acad Sci U S A 2001;98: 5116–21.
- Mathieu MC, Rouzier R, Llombart-Cussac A, et al. The poor responsiveness of infiltrating lobular breast carcinomas to neoadjuvant chemotherapy can be

explained by their biological profile. Eur J Cancer 2004;40:342-51.

- Rouzier R, Extra JM, Klijanienko J, et al. Incidence and prognostic significance of complete axillary downstaging after primary chemotherapy in breast cancer patients with T1 to T3 tumors and cytologically proven axillary metastatic lymph nodes. J Clin Oncol 2002;20:1304–10.
- Kuerer HM, Newman LA, Smith TL, et al. Clinical course of breast cancer patients with complete pathologic primary tumor and axillary lymph node response to doxorubicin-based neoadjuvant chemotherapy. J Clin Oncol 1999;17:460–9.
- 14. Gianni L, Zambetti M, Clark K, et al. Gene expression profiles of paraffin-embedded core biopsy tissue predict response to chemotherapy in patients with locally advanced breast cancer. ASCO Annual Meeting Proceedings 2004;22:501.
- **15.** Chang JC, Wooten EC, Tsimelzon A, et al. Gene expression profiling for the prediction of therapeutic response to docetaxel in patients with breast cancer. Lancet 2003;362:362–9.