Supervised Risk Predictor of Breast Cancer Based on Intrinsic Subtypes

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ABSTRACT

Purpose

To improve on current standards for breast cancer prognosis and prediction of chemotherapy benefit by developing a risk model that incorporates the gene expression–based “intrinsic” subtypes luminal A, luminal B, HER2-enriched, and basal-like.

Methods

A 50-gene subtype predictor was developed using microarray and quantitative reverse transcriptase polymerase chain reaction data from 189 prototype samples. Test sets from 761 patients (no systemic therapy) were evaluated for prognosis, and 133 patients were evaluated for prediction of pathologic complete response (pCR) to a taxane and anthracycline regimen.

Results

The intrinsic subtypes as discrete entities showed prognostic significance ($P = 2.26E-12$) and remained significant in multivariable analyses that incorporated standard parameters (estrogen receptor status, histologic grade, tumor size, and node status). A prognostic model for node-negative breast cancer was built using intrinsic subtype and clinical information. The C-index estimate for the combined model (subtype and tumor size) was a significant improvement on either the clinicopathologic model or subtype model alone. The intrinsic subtype model predicted neoadjuvant chemotherapy efficacy with a negative predictive value for pCR of 97%.

Conclusion

Diagnosis by intrinsic subtype adds significant prognostic and predictive information to standard parameters for patients with breast cancer. The prognostic properties of the continuous risk score will be of value for the management of node-negative breast cancers. The subtypes and risk score can also be used to assess the likelihood of efficacy from neoadjuvant chemotherapy.

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INTRODUCTION

Breast cancer is a heterogeneous disease with respect to molecular alterations, cellular composition, and clinical outcome. This diversity creates a challenge in developing tumor classifications that are clinically useful with respect to prognosis or prediction. Gene expression profiling by microarray has given us insight into the complexity of breast tumors and can be used to provide prognostic information beyond standard clinical assessment.1-7 For example, the 21-gene OncotypeDx assay (Genome Health Inc, Redwood City, CA) can be used to risk stratify early-stage estrogen receptor (ER)–positive breast cancer.4,5 Another strong predictor of outcome in ER-positive disease is proliferation or genomic grade.7-9 In addition, the 70-gene MammaPrint (Agenda, Huntington Beach, CA) microarray assay has shown prognostic significance in ER-positive and ER-negative early-stage node-negative breast cancer.2,3

The “intrinsic” subtypes luminal A (LumA), luminal B (LumB), HER2-enriched, basal-like, and normal-like have been extensively studied by microarray and hierarchical clustering analysis.1,6,10-12 Here, we study the utility of these subtypes alone and as part of a risk of relapse predictor in two cohorts: (1) patients receiving no adjuvant systemic therapy, and (2) patients undergoing paclitaxel, fluorouracil, doxorubicin, and cyclophosphamide (T/FAC) neo-adjuvant chemotherapy. The risk of relapse models were compared with standard models using pathologic stage, grade, and routine biomarker status (ER and HER2).
Samples and Clinical Data

Patient cohorts for training and test sets consisted of samples with data already in the public domain\textsuperscript{1,13-16} and fresh frozen and formalin-fixed paraffin-embedded (FFPE) tissues collected under institutional review board–approved protocols at the University of British Columbia (Vancouver, British Columbia, Canada), University of North Carolina (Chapel Hill, NC), Thomas Jefferson University (Philadelphia, PA), Washington University (St Louis, MO), and the University of Utah (Salt Lake City, UT). The training set for subtype prediction consisted of 189 breast tumor samples and 29 normal samples from heterogeneously treated patients given the standard of care dictated by their histology, stage, and clinical molecular marker status. The risk of relapse (ROR) models for prognosis in untreated patients were trained using the node-negative, untreated cohort of the Netherlands Cancer Institute (NKI) data set (n = 141).\textsuperscript{13} The subtype prediction and ROR models were independently tested for prognosis\textsuperscript{7,14,15} and chemotherapy response.\textsuperscript{16} The Hess et al.\textsuperscript{16} data set used for prediction of chemotherapy sensitivity was not associated with long-term outcome data and was evaluated based on information dictated by the manufacturer’s protocol (Qiagen RNeasy Midi Kit).

Nucleic Acid Extraction

Total RNA was purified from fresh-frozen samples for microarray using the Qiagen RNeasy Midi Kit according to the manufacturer’s protocol (Qiagen, Valencia, CA). The integrity of the RNA was determined using an Agilent 2100 Bioanalyzer (Agilent Technologies, Palo Alto, CA). The High Pure RNA Paraffin Kit (Roche Applied Science, Indianapolis, IN) was used to extract RNA from FFPE tissues (2 × 10-μm sections or 1.5-mm punches) for quantitative reverse transcriptase polymerase chain reaction (qRT-PCR). Contaminating DNA was removed using Turbo DNase (Ambion, Austin, TX). The yield of total RNA was assessed using the NanoDrop ND-1000 Spectrophotometer (NanoDrop Technologies Inc, Rockland, DE).

Reverse Transcription and qPCR

First-strand cDNA was synthesized using Superscript III reverse transcriptase (first Strand Kit; Invitrogen, Carlsbad, CA) and a mixture of random hexamers and gene-specific primers. PCR amplification and fluorescent melting curve analysis was done on the LightCycler 480 using SYBR Green I Master Mix (Roche Applied Science). A detailed protocol of the PCR conditions can be found in the Appendix (online only).

Microarray

Total RNA isolation, labeling, and hybridizations on Agilent human 1Av2 microarrays or custom-designed Agilent human 22k arrays were performed using the protocol described in Hu et al.\textsuperscript{14} All microarray data have been deposited into the Gene Expression Omnibus\textsuperscript{18} under the accession number GSE10886.

Identification of Prototypical Intrinsic Subtype Samples and Genes

To develop a clinical test that could make an intrinsic subtype diagnosis, we used a method to objectively select prototype samples for training and then predicted subtypes independent of clustering. To identify prototypic tumor samples, we started with an expanded “intrinsic” gene set comprised of genes found in four previous microarray studies.\textsuperscript{1,6,8,11} The normal-like class was represented using true “normals” from reduction mammoplasty or grossly uninvolved tissue, thus we have removed the normal-like class from all outcome analyses and consider this classification as a quality-control measure. A total of 189 breast tumors across 1,906 “intrinsic” genes were analyzed by hierarchical clustering (median centered by feature/gene, Pearson correlation, average linkage),\textsuperscript{17} and the sample dendrogram was analyzed using SigClust.\textsuperscript{20} A total of 122 breast cancers from 189 individuals profiled by qRT-PCR and microarray had significant clusters representing the “intrinsic” subtypes luminal A (LumA), luminal B (LumB), HER2-enriched, basal-like, and normal-like (Appendix Fig A1, online only). Four additional groups were identified in the training set as significantly reproducible clusters. All four of these groups have similar expression profiles as the luminal tumors and could represent intermediate states or tissue heterogeneity.

Gene Set Reduction Using Prototype Samples and Genes

A minimized gene set was derived from the prototypic samples using the qRT-PCR data for 161 genes that passed FFPE performance criteria established in Mullins et al.\textsuperscript{21} Several minimization methods were used, including top “N” test statistics for each group,\textsuperscript{22} top cluster index scores,\textsuperscript{23} and the remaining genes after “shrinkage” of modified t test statistics.\textsuperscript{24} Cross-validation (random 10% left out in each of 50 cycles) was used to assess the robustness of the minimized gene sets. The “N” t test method was chosen due to having the lowest cross-validation (random 10% left out of each iteration) error. The 50 genes selected and their contribution to distinguishing the different subtypes is provided in Appendix Figure A2 (online only).

Sample Subtype Prediction

The 50 gene set was compared for reproducibility of classification across three centroid-based prediction methods: Prediction Analysis of Microarray (PAM),\textsuperscript{24} a simple nearest centroid,\textsuperscript{6} and Classification of Nearest Centroid.\textsuperscript{25} In all cases, the subtype classification is assigned based on the nearest of the five centroids. Because of its reproducibility in subtype classification, the final algorithm consisted of centroids constructed as described for the PAM algorithm\textsuperscript{24} and distances calculated using Spearman’s rank correlation. The centroids of the training set using the 50-gene classifier (henceforth called PAM50) are shown in Appendix Figure A3 (online only).

### Table 1. Clinical Characteristics by Cohort

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Training Set</th>
<th>No Adjuvant Systemic Therapy</th>
<th>Neoadjuvant Chemotherapy</th>
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<tr>
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<tr>
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<tr>
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<td>33</td>
</tr>
<tr>
<td>Negative</td>
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<tr>
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<td>29</td>
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<td>Basal-like</td>
<td>56</td>
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<td>27</td>
</tr>
<tr>
<td>Normal-like</td>
<td>12</td>
<td>76</td>
<td>13</td>
</tr>
</tbody>
</table>

Abbreviations: RFS, relapse-free survival; NA, not applicable; ER, estrogen receptor.

\textsuperscript{*}Compiled from Ivshina et al,\textsuperscript{15} Loi et al,\textsuperscript{7} van de Vijver et al,\textsuperscript{13} Wang et al,\textsuperscript{14} Hess et al\textsuperscript{16} and University of North Carolina Microarray Database.\textsuperscript{17}
Prognostic and Predictive Models Using Clinical and Molecular Subtype Data

Univariate and multivariable analyses were used to determine the significance of the intrinsic subtypes (LumA, LumB, HER2-enriched, and basal-like) in untreated patients and in patients receiving neoadjuvant chemotherapy. For prognosis, subtypes were compared with standard clinical variables (tumor size \([T]\), node status \([N]\), ER status, and histologic grade), with time to relapse (ie, any event) as the end point. Subtypes were compared with grade and molecular markers (ER, progesterone receptor \([PR]\), HER2) for prediction in the neoadjuvant setting because pathologic staging is not applicable. Likelihood ratio tests were done to compare models of available clinical data, subtype data, and combined clinical and molecular variables. Categoric survival analyses were performed using a log-rank test and visualized with Kaplan-Meier plots.

Developing Risk Models With Clinical and Molecular Data

The subtype risk model was trained with a multivariable Cox model using Ridge regression fit to the node-negative, untreated subset of the van de Vijver cohort.\textsuperscript{13} A ROR score was assigned to each test case using correlation to the subtype alone (1) (ROR-S) or using subtype correlation along with tumor size (2) (ROR-C):

\[
\text{ROR-S} = 0.05 \cdot \text{basal} + 0.12 \cdot \text{HER2} + 0.34 \cdot \text{LumA} + 0.23 \cdot \text{LumB}
\]

\[
\text{ROR-C} = 0.05 \cdot \text{basal} + 0.11 \cdot \text{HER2} + 0.23 \cdot \text{LumA} + 0.09 \cdot \text{LumB} + 0.17 \cdot T
\]

The sum of the coefficients from the Cox model is the ROR score for each patient. To classify samples into specific risk groups, we chose thresholds from the training set that required no LumA sample to be in the high-risk group and no basal-like sample to be in the low-risk group. Thresholds were determined from the training set and remained unchanged when evaluating test cases. SiZer analysis was performed to characterize the relationship between the ROR score and relapse-free survival\textsuperscript{26} (Appendix Fig A4, online only). The 95% CIs for the ROR score are local versions of binomial CIs, with the local sample size computed from a Gaussian kernel density estimator based on the Sheather-Jones choice of window width.\textsuperscript{27}

Comparison of Relapse Prediction Models

Four models were compared for prediction of relapse: (1) a model of clinical variables alone (tumor size, grade, and ER status), (2) ROR-S, (3)
ROR-C, and (4) a model combining subtype, tumor size, and grade. The C-index was chosen to compare the strength of the various models. For each model, the C-index was estimated from 100 randomizations of the untreated cohort into two thirds training set and one thirds test set. The C-index was calculated for each test set to form the estimate of each model, and C-index estimates were compared across models using the two sample t-test.

## RESULTS

### Distribution of Intrinsic Subtypes in Comparison With Clinical Marker Status

Of the 626 ER-positive tumors analyzed in the microarray test set (Table 1), 73% were luminal (A or B), 11% were HER2-enriched, 5% were basal-like, and 12% were normal-like. Conversely, the ER-negative tumors comprised 11% luminal, 32% HER2-enriched, 50% basal-like, and 7% normal-like. The neoadjuvant study from Hess et al. provided an opportunity to analyze the subtype distribution.

Table 1 shows the intrinsic subtype distribution in ER-positive and ER-negative tumors. The intrinsic subtype distribution in ER-positive tumors is similar to that in ER-negative tumors, with a larger proportion of basal-like tumors in the ER-positive group. The proportion of luminal and HER2-enriched tumors is similar in both groups. The proportion of normal-like tumors is lower in the ER-positive group.

### Risk of Relapse Models for Prognosis in Node-Negative Breast Cancer

Cox models were tested using intrinsic subtype alone and together with clinical variables. Table 2 shows the multivariable analyses of these models in an independent cohort of untreated patients.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Model A</th>
<th></th>
<th>Model B</th>
<th></th>
<th>Model C</th>
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<tr>
<td></td>
<td>Hazard Ratio</td>
<td>P</td>
<td>Hazard Ratio</td>
<td>P</td>
<td>Hazard Ratio</td>
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<tr>
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<td>1.79</td>
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<td>&lt;.0001</td>
<td>2.90</td>
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<tr>
<td>Luminal B*</td>
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<td>&lt;.0001</td>
<td>2.88</td>
<td>&lt;.0001</td>
<td>2.54</td>
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<tr>
<td>ER status†</td>
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<td>.38</td>
<td>0.83</td>
<td>.34</td>
<td>0.83</td>
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<td>Tumor size‡</td>
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<td>Histologic grade¶</td>
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<table>
<thead>
<tr>
<th>Hazard Ratio</th>
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<tr>
<td>1.79</td>
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<tr>
<td>3.25</td>
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<td>1.36</td>
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<td>&lt;.0001</td>
</tr>
<tr>
<td>&lt;.0001</td>
<td>—</td>
<td>&lt;.0001</td>
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</tbody>
</table>

Abbreviation: ER, estrogen receptor.

*Luminal A class used as reference state in multivariable analysis.

†Hazard ratios for ER using positive marker in the numerator.

‡Size ≤ 2 cm versus ≥ 2 cm.

§Any positive node.

¶Grade encoded as an ordinal variable with three levels.

‖Significant P values indicate improved prediction relative to subtype alone.

*Significant P values indicate improved prediction relative to clinical data alone.
for identifying nonresponders to chemotherapy when using the ROR-S model to predict pCR (Fig 3A). The relationship between high-risk scores and a higher probability of pCR (Fig 3B) is consistent with the conclusion that indolent ER-positive tumors (LumA) are less responsive to chemotherapy. However, unlike ROR for prognosis, a plateau seems to be reached for the ROR versus probability of pCR, confirming the presence of significant chemotherapy resistance among the highest risk tumors.

**Subtype Prediction and Outcome on FFPE Samples Using qRT-PCR**

The subtype classifier and risk predictor were further validated using a heterogeneously treated cohort of 279 patients with FFPE samples archived between 1976 and 1995. The subtype classifications followed the same survival trends as seen in the microarray data, and the ROR score was significant for long-term relapse predictions (Appendix Fig A6A, online only). This old-age sample set was also scored for standard clinical markers (ER and HER2) by immunohistochemistry (IHC) and compared with the gene expression–based test. Analysis of ESR1 and ERBB2 by gene expression showed high sensitivity and specificity as compared with the IHC assay (Appendix Figs A6B and A6C). The advantages of using qRT-PCR versus IHC are that it is less subjective than visual interpretation and it is quantitative.

**DISCUSSION**

There have been numerous studies that have analyzed interactions between breast cancer intrinsic subtypes and prognosis, genetic alterations, and drug response. Because of the potential clinical value of subtype distinctions, we developed a standardized method of classification using a statistically derived gene and sample set that we have validated across multiple cohorts and platforms. The large and diverse test sets allowed us to evaluate the performance of the PAM50 assay at a population level and in relation to standard molecular
markers. An important finding from these analyses is that all of the intrinsic subtypes are present and clinically significant in terms of outcome predictions in cohorts of patients diagnosed with either ER-positive or ER-negative tumors (Fig 1). Thus the molecular subtypes are not simply another method of classification that reflects ER-positive or ER-negative tumors (Fig 1). The molecular subtypes are not simply another method of classification that reflects ER status.

Stratification of the subtypes within HER2clin-positive samples did not show significance in outcome predictions; however, there were fewer numbers and less follow-up in this category. Nevertheless, there was clear separation of the curves for those HER2clin-positive patients classified as HER2-enriched (worse prognosis) compared with those with luminal subtypes (better prognosis). We found that 6% of HER2clin-positive tumors were classified as basal-like. It has been suggested that HER2clin-positive tumors expressing basal markers may have worse outcome when given a chemotherapeutic regimen of trastuzumab and vinorelbine.31

Approximately one third of the HER2-enriched expression subtype were not HER2clin-positive tumors, suggesting the presence of an ER-negative, nonbasal subtype that is not driven by HER2 gene amplification. The prototype samples selected to represent the HER2-enriched group had high expression of the 17q12-21 amplicon genes (HER2/ERBB2 and GRB7, FGFR4 (5q35), TMEM45B (11q24), and GPR160 (3q26). In addition, other growth factor receptors such as epidermal growth factor receptor are included within the PAM50 and could potentially also contribute to the HER2-enriched genomic classification.

We found that approximately 10% of breast cancers were classified as normal-like and can be either ER-positive or ER-negative and have an intermediate prognosis. Because the normal-like classification was developed by training on normal breast tissue, we suspect that the normal-like class is mainly an artifact of having a high percentage of normal “contamination” in the tumor specimen. Other explanations include a group of slow-growing basal-like tumors that lack expression of the proliferation genes or a potential new subtype that has been referred to as claudin-low tumors.32 Detailed histologic, immunohistochemical, and additional gene expression analyses of these cases are needed to resolve these issues. Because of the uncertainties, however, the normal-like samples were removed when modeling ROR.

The multivariable analysis for prognosis (ie, no systemic treatment) suggested that the best model was to use subtype with pathologic staging. Because pathologic staging is not available at diagnosis in the neoadjuvant setting, we used histologic grade and clinical biomarkers as the standard for prediction of chemotherapy response before resection. In this context, only the subtypes LumB and basal-like were predictive in the multivariable analysis that included histologic grade, ER, PR, and HER2 status (note that the Hess et al16 study did not incorporate trastuzumab into the regimen). The ROR score from the subtype-alone model was also the most predictive of neoadjuvant.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Model A Odds Ratio</th>
<th>P</th>
<th>Model B Odds Ratio</th>
<th>P</th>
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<tbody>
<tr>
<td>Basal-like†</td>
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<td>.021</td>
<td>3.06</td>
<td>.016</td>
</tr>
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<td>.133</td>
<td>1.99</td>
<td>.117</td>
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<tr>
<td>Luminal B†</td>
<td>2.42</td>
<td>.036</td>
<td>2.43</td>
<td>.035</td>
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<td>ER status†</td>
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<td>.133</td>
<td>1.31</td>
<td>.124</td>
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Abbreviations: ER, estrogen receptor; PR, progesterone receptor. *Luminal A class used as reference state in multivariable analysis. †Hazard ratios for ER, PR, and HER2 are positive marker in the numerator. ‡Grade encoded as an ordinal variable with three levels. §Significant P values indicate improved prediction relative to subtype alone. ¶Significant P values indicate improved prediction relative to clinical data alone.
response. One of the major benefits of the ROR predictor is the identification of patients in the LumA group who are at a low ROR on the basis of pure prognosis and for whom the benefit from neoadjuvant therapy is unlikely. Thus the ROR predictor based on subtypes provides similar information as the OncotypeDx Recurrence Score for patients with ER-positive, node-negative disease.5,6 Furthermore, the PAM50 assay provides a ROR score for all patients, including those with ER-negative disease, and is highly predictive of neoadjuvant response when considering all patients.

In summary, the intrinsic subtype and risk predictors based on the PAM50 gene set added significant prognostic and predictive value to pathologic staging, histologic grade, and standard clinical molecular markers. The qRT-PCR assay can be performed using archived breast tissues, which will be useful for retrospective studies and prospective clinical trials.

AUTHORS’ DISCLOSURES OF POTENTIAL CONFLICTS OF INTEREST

Although all authors completed the disclosure, the following author(s) indicated a financial or other interest that is relevant to the subject matter under consideration in this article. Certain relationships marked with a “C” were compensated. For a detailed description of the disclosure categories, or for more information about ASCO’s conflict of interest policy, please refer to the Author Disclosure Declaration and the Disclosures of Potential Conflicts of Interest section in Information for Contributors.

REFERENCES


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Appendix

Reverse Transcription and Real-Time Quantitative Polymerase Chain Reaction

First-strand cDNA was synthesized from 1.2 μg of total RNA using Superscript III reverse transcriptase (first Strand Kit; Invitrogen, Carlsbad, CA) and a mixture of random hexamers and gene-specific primers. The reaction was held at 55°C for 60 minutes and then 70°C for 15 minutes. The cDNA was washed on a QIAquick polymerase chain reaction (PCR) purification column (Qiagen Inc, Valencia, CA) and stored at −80°C in 25 mmol/L of Tris and 1 mmol/L of EDTA until further use. Each 5-μL PCR reaction included 1.25 ng (0.625 ng/μL) of cDNA from samples of interest or 10 ng (5 ng/μL) for reference, 2 pmol of both upstream and downstream primers, and LightCycler 480 SYBR Green I Master Mix (Roche Applied Science, Indianapolis, IN). Each run contained a single gene profiled in duplicate for test samples, reference sample, and negative control. The reference sample cDNA comprised an equal contribution of Human Reference Total RNA (Stratagene, La Jolla, CA) and the breast cell lines MCF7, ME16C, and SKBR3. PCR amplification was performed with the LightCycler 480 (Roche Applied Science, Indianapolis, IN) using an initial denaturation step (95°C, 8 minutes) followed by 45 cycles of denaturation (95°C, 4 seconds), annealing (56°C, 6 seconds with 2.5°C/s transition), and extension (72°C, 6 seconds with 2°C/sec transition). Fluorescence (530 nm) from the dsDNA dye SYBR Green I was acquired each cycle after the extension step. The specificity of the PCR was determined by postamplification melting curve analysis: samples were cooled to 65°C and slowly heated at 2°C/s to 99°C while continuously monitoring fluorescence (10 acquisitions/1°C).
Hierarchical clustering and SigClust analysis of microarray data using 1,906 "intrinsic" genes and 189 samples. The SigClust algorithm statistically identifies significant/unique groups by testing the null hypothesis that a group of samples is from a single cluster, where a cluster is characterized as a multivariable normal distribution. SigClust was run at each node of the dendrogram beginning at the root and stopping when the test was no longer significant ($P > .001$). Statistical selection using SigClust identified nine significant groups, including the previously identified subtypes designated as luminal A (dark blue), luminal B (light blue), HER2-enriched (pink), normal-like (green), and basal-like (red).
Fig A2. Focused heatmap of Classification by Nearest Centroids (ClaNC) selected genes for each subtype. The ClaNC algorithm was optimized to select 10 genes per class for a total of 50 genes. The 10 genes for each class are shown as red/green according to their expression in a class. Black indicates that gene was not selected for the given class.
Fig A3. Heatmap of the centroid models of subtype. The centroids were constructed using the Classification by Nearest Centroids selected genes and calculated as described for the Prediction Analysis of Microarray algorithm. The expression values are shown as red/green according to their relative expression level.
Fig A4. Developing a continuous risk score based on subtypes and clinical variables. (A) A family of smoothing functions illustrates the general linear relationship between the risk of relapse (ROR) score and relapse-free survival at 5 years. (B) Significance of the slopes in the smoothed lines is plotted for each bandwidth across the range of scores. Blue indicates a significant positive slope, purple indicates nonsignificant slope, and green indicates too few data for inference.

Fig A5. Risk classification for training set of untreated patients using a combined model of intrinsic subtypes and clinical variables. (A) Risk of relapse based on the combined model with low-risk scores less than 29 (black), moderate-risk scores between 29 and 53 (green), and high-risk scores ≥ 53 (red). (B) Kaplan-Meier plot and significance of the risk score shown for 141 training set cases. LumA, luminal A; LumB, luminal B.
Fig A6. Analysis of an old-aged formalin-fixed, paraffin-embedded patient cohort. (A) The combined risk of relapse (ROR)-C model predicting high, medium, and low risk of relapse in 279 patients from the University of British Columbia. (B) Receiver operating characteristic curve analysis of quantitative reverse transcriptase polymerase chain reaction (qRT-PCR) data for ESR1 versus immunohistochemistry (IHC) data. (C) Receiver operating characteristic curve analysis for qRT-PCR data for HER2/ERBB2 versus HER2clin-positive (IHC and/or fluorescent in situ hybridization). PPV, positive predictive value; NPV, negative predictive value; AUC, area under the curve.
Introducing Molecular Subtyping of Breast Cancer Into the Clinic?

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Breast cancer is a collection of diseases demonstrating heterogeneity at the molecular, histopathologic, and clinical level. The diversity at the molecular level is manifested in differences in gene expression patterns (both of mRNA and microRNA), different frequencies and magnitudes of genomic aberrations, and differential protein expression across breast tumors, even among those of similar histopathologic type.\(^1\)\(^2\) In addition to this, and influencing this diversity in tumor cells, is the diversity in the microenvironment, reflecting different degrees of involvement of various biologic processes.\(^3\)\(^-\)\(^5\) The molecular heterogeneity is reflected in the clinical course of the diseases and responses to treatment.

During the last 10 years, whole-genome analyses using microarrays have revolutionized cancer research. Such studies of breast tumors have led to the identification of five molecular subtypes associated with differences in patient survival,\(^6\)\(^-\)\(^7\) as well as numerous gene expression signatures associated with different clinical parameters.\(^8\)\(^-\)\(^14\) Some of these have formed the basis for commercialized tests that are now being assessed in large clinical trials;\(^15\) however, none has taken into account the heterogeneity represented by the molecular subtypes.

In this issue of Journal of Clinical Oncology, Parker et al\(^16\) report on a risk prediction model for breast cancer developed from expression data of 50 genes representing the five intrinsic molecular subtypes. They show that the intrinsic subtypes are present in several tumor cohorts, both those positive or negative for estrogen receptor (ER) and human epidermal growth factor receptor 2 (HER-2), and are associated with significant differences in relapse-free survival in several published breast cancer microarray data sets. Importantly, this illustrates that the molecular subtypes are not just recapitulating the classic clinical markers for classification; for example, basal-like tumors cannot simply be substituted by triple-negative (ER negative/herceptin negative) status. In addition, they show in a multivariable analysis that the subtypes are able to predict nonresponse to a neoadjuvant paclitaxel plus fluorouracil, doxorubicin, and cyclophosphamide regimen with high sensitivity and high negative predictive value, although at the expense of specificity. The predictor was built from a microarray training set consisting of a published subcohort of patients with node-negative untreated breast cancer. A model that incorporated both the intrinsic subtypes and tumor size resulted in improved risk prediction for relapse in untreated patient cohorts compared with either subtypes or clinical markers alone. A final validation step by quantitative reverse transcriptase polymerase chain reaction test using formalin-fixed paraffin-embedded material showed similar subtype distribution and prognostic capability of the predictor. A strength of the study is the ability to assign subtypes with a limited gene set using archived tumor tissue with extensive follow-up information.

The predictor classified patients into three risk groups (high, medium, and low), two of which contained each of the subtypes; the low-risk class consisted only of luminal A type tumors. Conversely, the risk predictor divided patients with luminal A type tumors into both low- and intermediate-risk groups, which is of significant value if this may help to identify patients who might be spared aggressive treatment. Although patients with luminal A type tumors are associated with relatively good prognosis, a significant portion experience relapse, indicating heterogeneity within this group.\(^17\) It remains to be seen if this three-category risk predictor is superior to any of the other published prognostic predictors based on gene expression signatures. Another strength of this model is that it provides prognostic value for all types of breast cancer patients, irrespective of ER and node status as well as disease stage. Whether this same model may robustly predict nonresponse to treatment is still unclear, given that this study predicted residual disease in a preoperative cohort designed to predict pathologic complete response, and in which patients with residual disease comprised the major part.\(^18\)

These five molecular subtypes, which were validated extensively, are profoundly different in their gene expression patterns (including microRNA expression), genetic alterations, and distribution of mutations and normal variants.\(^19\)\(^-\)\(^27\) This suggests that they are developing along distinct pathways and are different mechanistically. This is especially true for the luminal A and basal-like subtypes. These are inversely correlated and protrude in any genome-wide study at the genomic, transcriptomic, and proteomic level. They probably originate from different lineages of progenitor cells (or stem cells) at different stages of differentiation. The distinctiveness of the three other subtypes is more unclear. The ERBB2-positive group is characterized by strong expression of the ERBB2 oncogene and a few other genes located in the same region on chromosome 17q as a result of amplification. However, amplification of this region is also seen in luminal B tumors, a subtype of tumors expressing ER and other estrogen-related genes, but which also share some expression characteristics with basal-like tumors. The normal-like tumor subtype, originally defined by its similarity in expression with normal tissue and benign tumors, is not just a reflection of poor tissue sampling, but is nevertheless not...
associated with a clear signature. Hence, these latter groups may reflect heterogeneity of the two main separate breast cancer types, luminal and basal-like. Genomic rearrangements and genetic instability occur at different stages in tumor development and bring about some of the major differences seen among the subtypes. In light of the cancer stem-cell concept, the different subtypes may develop along the differentiation from a multipotent cancer stem cell to the three different cell populations in the breast. The heterogeneity observed may result from specific alterations in genes and pathways on a luminal or basal background, which, together with interactions with the microenvironment, result in these prodigiously different phenotypes. This work emphasizes the clinical value of subtype stratification, which is important for discovering biomarkers that may act differently within the context of these groups. Additional research in this field will move from the descriptive approaches to a more hypothesis-driven and mechanistically driven strategy to identify the drivers of the coordinate expression of genes that comprise the molecular subtypes.

Getting closer to a complete understanding of the complexity of the molecular aberrations underlying breast cancer initiation and progression, and the impact this has on treatment strategies, can only be achieved by integrated analytic approaches. Realizing the existence of several molecular subtypes of breast cancer, perhaps genuinely different in the cell of origin, and designing research protocols and clinical studies accordingly, will bring us closer to successful treatment of breast cancer patients. It is hoped that a classification scheme that identifies more homogeneous tumors built on the intrinsic molecular subtypes will lead in this direction—moving further away from the “one-size-fits-all” concept of therapy and toward personalized treatment.

AUTHOR’S DISCLOSURES OF POTENTIAL CONFLICTS OF INTEREST

The author(s) indicated no potential conflicts of interest.

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