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Gene expression signature in organized and growth arrested mammary acini predicts good outcome in breast cancer

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Running title: A mammary gland differentiation signature predictive of good breast cancer outcome
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Key words: microarray, 3D culture, extracellular matrix, gene expression profile, breast cancer.
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ABSTRACT:

To understand how non-malignant human mammary epithelial cells (HMEC) transit from a disorganized proliferating to an organized growth arrested state, and to relate this process to the changes that occur in breast cancer, we studied gene expression changes in non-malignant HMEC grown in three-dimensional cultures, and in a previously published panel of microarray data for 295 breast cancer samples. We hypothesized that the gene expression pattern of organized and growth arrested mammary acini would share similarities with breast tumors with good prognoses. Using Affymetrix HG-U133A microarrays, we analyzed the expression of 22,283 gene transcripts in two HMEC cell lines, 184 (finite life span) and HMT3522 S1 (immortal non-malignant), on successive days post-seeding in a laminin-rich extracellular matrix assay. Both HMECs underwent growth arrest in G0/G1 and differentiated into polarized acini between days 5 and 7. We identified gene expression changes with the same temporal pattern in both lines. We show that genes that are significantly lower in the organized, growth arrested HMEC than in their proliferating counterparts can be used to classify breast cancer patients into poor and good prognosis groups with high accuracy. This study represents a novel unsupervised approach to identifying breast cancer markers that may be of use clinically.
INTRODUCTION:

Loss of growth control and disruption of tissue architecture are among the earliest hallmarks of cancer. We hypothesized that the gene expression changes that occur during the organization and growth arrest of cultured mammary acini would share similarities with breast tumors that had a good prognosis. To test this hypothesis, we examined temporal changes in gene expression in two non-malignant human mammary epithelial cells (HMEC) grown in a three-dimensional laminin-rich extracellular matrix (3D lrECM) assay (1). In this model, single non-malignant breast epithelial cells form polarized growth arrested multicellular structures that resemble acini over a period of several days. This transition from the unpolarized actively dividing state to a polarized non-dividing state is the reverse of what happens in the early stages of tumorigenesis.

We used two non-malignant HMEC, a non-immortalized HMEC strain, 184 (2, 3), and spontaneously immortalized cell line, HMT3522-S1 (4), and monitored the changes in gene and protein expression as the cells formed acinar structures in 3D lrECM cultures. During the process of self-organization and withdrawal from the cell cycle, we noted a progressive hypophosphorylation of the retinoblastoma gene product (Rb) and an induction of the cyclin-dependent kinase (cdk) inhibitor p27kip1. Analysis of the changes in gene expression in both cell lines allowed the identification of sets of commonly regulated genes.

While established breast cancer prognostic markers such as tumor size, grade, lymph node and hormone receptor status are useful in predicting survival in large populations (5-7), there is a pressing need to develop better prognostic signatures to predict recurrence and overall survival. A particular benefit would be the identification of
patients with good prognoses whose tumors are highly unlikely to recur and who nevertheless are being treated with cytotoxic chemotherapies (8). The advent of gene expression technologies has greatly aided the identification of molecular signatures with value for tumor classification and prognosis prediction (9-14). Van de Vijver et al. have developed a 70-gene signature that effectively stratifies patients into good and poor prognosis groups (15, 16). Paik et al. have proposed a 21-gene signature with which to calculate a “recurrence score” that predicts the likelihood of recurrence in estrogen receptor (ER)-positive lymph node-negative patients (17). In each of these studies, the predictive signatures have been derived by using a training set of patients of known outcome, followed by testing these signatures in a validation set of patients. In contrast, our approach has been to identify directly the genes whose expression changes as cultured mammary epithelial cells transition from a disorganized to an organized state, and to then test as proof of principle the possible utility of these genes as prognostic markers in a validation set of patients.

MATERIALS AND METHODS:

Cell Culture.

Finite-lifespan 184 HMEC were obtained from reduction mammoplasty tissue and grown in a serum-free MCDB 170 medium (MEGM; Clonetics Division of BioWhittaker, Walkersville, MD, USA), as described previously (2, 18). HMT-3522 S1 mammary epithelial cells were cultured in H14 medium (DMEM/F12 containing 250 ng/ml insulin, 10 µg/ml transferrin, 2.6 ng/ml sodium selenite, $10^{-10}$ M estradiol, $1.4 \times 10^{-6}$ M hydrocortisone, 10 ng/ml EGF and 5 µg /ml prolactin). The cells were cultured in a
3D laminin-rich extracellular matrix (Matrigel, BD Biosciences), as described (19). The colonies were isolated from the Matrigel in ice-cold PBS/5 mM EDTA after 3, 5, 7, and 10 days. For western blot analysis, the colonies were lysed in 150 mM NaCl, 1 % NP-40, 50 mM Tris, pH 8.0. RNA was isolated using the RNeasy kit (Qiagen), according to the manufacturer’s instructions.

Indirect immunofluorescence and image acquisition.

Acinar structures were fixed on glass slides in methanol-acetone (1:1) at -20°C for 10 minutes and air dried. A primary block was performed in an IF Buffer (130 mM NaCl, 7 mM Na2HPO4, 3.5 mM NaH2PO4, 7.7 mM NaN3, 0.1% bovine serum albumin, 0.2% Triton X-100, 0.05% Tween-20) + 10% goat serum for 1 hour at room temperature. A secondary block was performed in IF Buffer + 10% goat serum + 20µg/ml goat anti-mouse F(ab’)2 fragment for 30 minutes. The primary antibody, rat anti-alpha-6 integrin (Chemicon International), was diluted 1:50 in the latter blocking buffer and incubated overnight (15-18 hours) at 4°C. Secondary antibodies conjugated with fluorescent dye FITC or Texas Red were diluted 1:100 in blocking buffer and incubated 1 hour at room temperature. Immunofluorescent images were acquired using an inverted microscope equipped with a digital camera and SPOT software. Confocal analysis was performed using a Zeiss 410 confocal microscopy system. The images presented are representative of two or more independent experiments. All the images were converted to TIFF format and arranged using Adobe Photoshop 5.0.

Flow cytometry.

To analyze the DNA content, acini were suspended in 300 µl 0.25% trypsin and incubated at 37°C for exactly 10 minutes. The dispersed cells were washed three times in
PBS and fixed in 40% ethanol at 4°C overnight. They were then incubated with 500 µg/ml RNaseA in PBS at 37°C for 30 minutes and stained with 69µM propidium iodide solution in PBS at room temperature for 30 minutes. The DNA content was determined by flow cytometry using a FACScan (Becton Dickinson, Hialeah, FL), and the data were analyzed with Cell Quest software (Becton Dickinson).

Immunoblotting.

Whole-cell lysates (20–100 µg protein) were separated by SDS-PAGE and transferred electrophoretically to PVDF membranes (Millipore, Billerica, MA). After blocking in 10% nonfat dry milk for 90 minutes at room temperature, they were incubated with the following antibodies at 1:500 to1:1000 dilution in 2% nonfat dry milk for 2 hours at room temperature or overnight at 4°C: rabbit anti-phosphorylated Rb at S807/811 and S795 (Cell Signaling Technology, Beverly, MA), rabbit anti-phosphorylated Rb at S612 and T821 (Biosource, Camarillo, CA), and mouse anti-p27 kip1 (BD Transduction Laboratories). The blots were then incubated with HRP-conjugated sheep anti-mouse IgG or anti-rabbit IgG at 1:2000 in 2% nonfat dry milk for 90 minutes at room temperature. The blots were developed using Super Signal West Pico Chemiluminescent Substrate (Pierce, Rockford, IL).

Survival analysis.

A database consisting of the microarray profiles of 295 human breast tumors with the associated clinical data (15) was obtained from Rosetta Inpharmatics (http://www.rii.com/publications/2002/nejm.html). For survival analysis of the 19 individual marker genes, the patients were stratified into quartiles for expression of each marker, and the survival curves were computed using the method of Kaplan and Meier.
Statistical significance was determined using the log-rank test. Statistical analyses were performed using Graphpad Prism. For survival analysis of the set of 249 marker genes, the patients were stratified into two groups using GeneSpring software by hierarchical cluster analysis with a distance metric of the expression pattern of all 249 genes. Kaplan-Meier survival curves, log-rank statistics, and the estimated hazard ratio for these two groups were computed using the Excel add-in EcStat.

**Microarray hybridization and analysis.**

Cell samples were harvested in duplicate at three time points, 3, 5, and 7 days, after seeding in lrECM. Purified total cellular RNA was biotin-labeled and hybridized to human oligonucleotide microarrays (Affymetrix HG-U133A), as previously described (20). Experiments with Affymetrix-present P-call rates of >30% were included in the analysis. Signal values from each of the 22,283 probe sets were calculated by means of robust multi-array analysis (RMA) (21) using Bioconductor ([http://www.bioconductor.org](http://www.bioconductor.org)) in the R computing environment. The signal values were inverse log2 transformed and then imported into GeneSpring software (SiliconGenetics); and each array was normalized to its median signal intensity. The genes were normalized to the mean of the 3-day time point for each cell type independently. For Method 1 of selecting significantly differential temporally co-regulated genes (Figure 2): Significantly up-regulated genes in each cell specimen were identified by first selecting the genes induced at least 1.5 fold in at least one of the six conditions and then performing an ANOVA analysis as a function of time. Variances were calculated using the cross-gene error model (GeneSpring), p-value cutoff 0.05, multiple testing correction: Benjamini and Hochberg False Discovery Rate. About 5% of the identified genes in each set would be
expected to pass this restriction by chance. Significantly down-regulated genes were identified in the same manner after normalizing to the 7-day time point. Genes that were up- or down-regulated early in each cell line were selected from the significantly up- or down-regulated gene lists. The early genes were defined as those with a mean expression at 5 days of at least 50% of their mean expression at 7 days. For Method 2 of selecting significantly differential, temporally co-regulated genes (Figure 6A): We first selected all genes that were at least 1.5 fold differential (up or down) in at least one of the four samples from the two later time points (days 5 and 7) for either cell specimen. We then performed an ANOVA as a function of time. Variances were not assumed equal (Welch ANOVA), p-value cutoff 0.05, multiple testing correction: Benjamini and Hochberg False Discovery Rate. From this list of genes that were significantly differential in either cell line, we then identified genes that were up- or down-regulated early (mean expression at 5 days of at least 50% of their mean expression at 7 days) or late in each cell line. We then identified those genes that were coordinately regulated in both cell lines.

RESULTS:

Temporal analysis of gene expression in two non-malignant human breast epithelial cells grown in 3D lrECM cultures.

To identify consistent changes in gene and regulatory protein expression levels, we characterized two independently derived non-malignant HMECs: one finite life-span strain (184) (2) and one spontaneously immortalized line (HMT3522 S1) (4). Both cells formed acinus-like structures with similar morphology and basal polarity when cultured
from single cells in lrECM (Figure 1A and 1B). We performed temporal studies to
determine when, and in what order, changes in critical cell cycle regulatory molecules
occurred in non-malignant HMECs in 3D lrECM cultures. Flow cytometric analyses
indicated that, after undergoing a limited number of cell divisions, the majority of non-
malignant S1 cells accumulated in the G0/G1 phase of the cell cycle by day 7 (Figure
1C). The product of the retinoblastoma susceptibility gene (Rb), a central player in the
G1/S transition, is inactivated by phosphorylation, allowing cell cycle progression. Rb
inactivation occurs through the sequential actions of cyclin D/cdk4 and 6, and of cyclin
E-ckd2 complexes (22). We analyzed cell cycle regulators known to affect G1
checkpoints in a time-dependent manner in S1 cells in 3D cultures. Phosphorylation of
several sites on Rb was found to gradually decrease between days 5 and 10 in these non-
malignant cells, consistent with the growth-suppressive role of the hypophosphorylated
form (Figure 1D). Cyclins D1, E, and A, as well as their binding partners—cdks 4, 6,
and 2—also decreased during this period (data not shown). In contrast, the cdk inhibitor
p27kip1 increased between days 5 and 10 (Figure 1D). The pronounced down-modulation
of Rb phosphorylation and the elevation of p27kip1 protein levels were changes observed
in both the S1 and 184 cells (data not shown). Although other studies have used different
HMEC (MCF10A) to determine how cells can escape normal growth control (23-26), the
mechanisms by which mammary cells actually initiate and maintain growth arrest during
the process of acini formation in the context of 3D-lrECM remain to be determined.

Global gene expression analysis of the time course of HMECs in 3D.

To probe systematically the molecular changes that accompany acinus formation,
we analyzed the expression profiles of 22,283 transcripts using Affymetrix HG-U133A
microarrays. Microarray experiments were performed with biological duplicates using RNA samples harvested from S1 and 184 cells, after 3, 5, and 7 days’ culture in lrECM. Since growth arrest and polarization occurred with similar kinetics in both cell types (Figure 3A), we reasoned that the gene expression changes important for these processes would follow a common temporal pattern in both cell lines, and that changes that were cell type-specific could be disregarded.

We first identified genes that showed at least 1.5-fold changes during the time course in the individual cell specimens (ANOVA, p < 0.05) (within this window, 363 genes were up-regulated and 117 genes down-regulated in 184 cells; 234 genes were up-regulated and 351 genes down-regulated in S1 cells). We then divided these lists into genes whose expression changed ‘early’ by our definition (between days 3 and 5) or ‘late’ (between days 5 and 7) in S1 and 184. Finally, we identified the genes from each temporal group that were common to both cell types (Figure 2). A total of 60 genes with common temporal patterns were identified, including 21 genes that were up-regulated early, 11 genes that were up-regulated late, 6 genes that were down-regulated early, and 22 genes that were down-regulated late (Figure 3B and 3C and supplementary material).

Correlation of the differentially expressed genes with survival of breast cancer patients.

To relate the process of acinar development in 3D lrECM cell cultures to the changes that occur in breast cancer, we examined the expression levels of the differentially regulated genes identified using our model using previously published microarray data for a panel of 295 breast cancer samples from the fresh-frozen-tissue bank of the Netherlands Cancer Institute, including 151 lymph node-negative and 144
lymph node-positive patient samples (15). Fifty five of the 60 genes selected in our 3D culture analysis were included on these microarrays. We looked at 5- and 10-year survival data and applied Student’s t-test to determine how many of the genes modulated in 3D cultures showed survival-associated expression changes. T-tests were performed to determine whether the difference in the expression level of a given gene in two groups (e.g., patients who survived five years versus patients who did not) was large enough that it was not likely to be due to chance. The numbers and percentages of genes exhibiting significantly different expression in the tumors of patients with differential survival (p < 0.05) were tabulated for a) all the genes represented on the microarrays, b) genes selected on the basis of differential expression during the 3D lrECM timecourse, or c) randomly generated gene lists (Table 1). The percentage of genes with survival-associated expression changes was highest for those genes down-regulated late (between days 5 and 7) in the time course. The percentage for this gene list exceeded those of the unfiltered list of all 25,773 genes represented on the arrays, 5 random gene lists, and all other 3D lrECM gene lists. The list of genes that were down-modulated late in the lrECM timecourse showed a marked enrichment in genes whose expression level correlated to 5 (68%) and 10 years survival (53%). The levels of the majority of the late down-regulated genes were higher in patients who died within 5 or 10 years (ACTB, VRK1, ODC1, CKS2, FLJ10036, FLJ10540, FOXM1, RRM2, TRIP13, CDKN3, STK6, FLJ10517, TUBG1, ACTN1, TNFRSF6B and EPH2), while the levels of three genes (DUSP4, HBP17 and EIF4A1) were lower in these patients.

We identified 22 genes in all that were down-regulated in both HMECs between days 5 and 7 of lrECM culture. Of these, 19 genes were represented in the published data
set for the 295 patient tumor samples. We stratified the 295 tumors into quartiles based on the relative expression level of each of the genes in the selected set, and further analyzed the relationship of the expression level of each individual gene to survival (Figure 4). The resulting Kaplan-Meier curves showed that gene expression levels correlated significantly with outcome for 14 of the 19 selected markers. For 13 of the 14 markers, gene expression was lower in tumors from patients with better outcomes, while in one case (DUSP4) gene expression was lower in tumors from patients with poorer outcomes.

Collective gene signatures have the potential to discriminate among clinical endpoints more accurately than markers used individually. Hence, we tested the ability of our set of 19 genes to classify breast cancer patients into prognostic groups. We used hierarchical cluster analysis to separate the patients into groups and then determined the overall 10-year survival rates for these groups (Figure 5). The cluster analysis separated the patients into five groups, three of which had tumors that expressed comparatively lower levels of most of the 19 genes, and two of which expressed higher levels. The 10-year survival rates for these 5 groups were 95, 84, 67, 61, and 54% respectively.

To test whether other sets of genes down-regulated late in the IrECM timecourse identified by using other selection strategies would also include useful breast cancer markers, we applied a second selection strategy (Figure 6A) and tested the ability of the resulting gene set to predict breast cancer prognosis. This second method was less restrictive than the first, and resulted in the identification of 287 genes that were significantly down-regulated late in the 3D time course of both HMEC specimens (for complete gene list and gene expression information, see supplemental information).
Seventeen of the 22 genes selected using Method 1 were also included in the 287 genes selected using Method 2. We tested the ability of these genes to predict breast cancer prognosis by using hierarchical cluster analysis in the same set of previously published microarray data from 295 breast cancers (15). A large majority, 249 of the 287 genes were included on these microarrays. Of the 17 overlapping (methods 1 and 2) genes on the Affymetrix chips, 15 were present on the Rosetta chips: ACTB, ACTN1, CDK3, CKS2, DUSP4, EPHA2, HBP17, FOXM1, ODC1, RRM2, STK6, TNFRSF6B, TRIP13, TUBG1, VRK1.

Hierarchical cluster analysis using the 249 gene signature classified the samples into two groups of approximately equal numbers of tumors (Figure 6B). Overall 10 year survival rates were 90% (138 of 154) for the good prognosis group and 59% (83 of 141) for the poor prognosis group. To assess the significance of these predictions and take into account patients that could not be followed the entire length of the study, we performed a Kaplan-Meier analysis. The results show that the 249 gene profile was highly informative in identifying patients with poor outcome (log rank p = 2.7x10^{-10}) (Figure 6C). The estimated hazard ratio for poor outcome (failure to survive) in the group with the poor prognosis signature as compared with the good prognosis signature was 4.7 [95% confidence interval 2.8 – 7.9].

DISCUSSION:

Three-dimensional (3D) laminin-rich (lrECM) cultures permit non-malignant cells to exhibit self-organizing properties. Such cultures provide models that allow the
study of processes that are aberrant in breast cancer (1, 19, 27, 28). To understand how breast epithelial cells transition from a proliferating, unorganized state to a resting, organized state, and to relate this process to the opposing changes that occur in clinical breast cancer, we performed genome-wide gene expression profiling for two non-malignant HMECs in 3D cultures, and utilized published data from a panel of 295 breast cancer samples. We show that genes that are down-regulated as HMEC transition from a proliferative to an acinar phenotype can be used collectively as signatures to predict clinical breast cancer prognosis.

Our approach represents a new way to identify genome-wide cancer prognostic markers. We based all the marker selection steps on HMEC cultured in 3D lrECM. The 3D model system provided a means to focus on epithelial cells themselves, and a defined, highly relevant biological process - the formation of breast acini. Whereas the stroma is absent, the 3D lrECM assays appears to substitute for myoepithelial cells and other signals that are needed to form an organized acini (29). Differentially expressed genes identified using this model system are likely to be functionally linked to the transformation-relevant process. Further, we have applied an unsupervised method (hierarchical cluster analysis) to classify the patient samples using selected markers. Hence, neither our method of marker selection nor our sample classification method relies on any clinical information.

Gene-expression profiling of tumors using DNA microarrays is a promising method for predicting prognosis and treatment response in cancer patients (17, 30-32). Two studies have recently employed genome-wide microarray analysis to identify gene signatures that predict prognosis in breast cancer. The profiles studied by both groups of
researchers were reported to be more powerful predictors of the outcome of disease than standard systems based on clinical and histological criteria. The study by van’t Veer et al. (16) used the supervised classification of a primary data set of 78 tumor samples to identify a 70-gene signature that divided the samples into classes of poor and good prognosis. Validation of the classifier by a second overlapping set of 295 tumors showed that it accurately predicted 10-year survival in breast cancer patients (15). Overall, the 10-year survival rates were 54.6% ±4.4% and 94.5% ±2.6% for the poor and good prognosis groups, respectively. Their estimated hazard ratio for poor outcome (distant metastases) in the group with a poor-prognosis signature, as compared with the group with the good-prognosis signature, was 5.1 (95% confidence interval, 2.9 to 9.0; \( P<0.001 \)). In a similar study, Wang and collaborators (33) also used supervised classification in a training set of 115 breast tumor samples. They identified a 76-gene signature, which was verified in an independent set of 171 breast tumors. Their estimated hazard ratio for poor outcome (distant metastases within 5 years) in the group with the poor prognosis signature as compared with the group with the good prognosis signature was 5.7 (95% confidence interval, 2.6 to 12.4). Our 249-gene signature predicted 10-year survival rates of 59% and 90% for poor and good prognosis groups, respectively. The estimated hazard ratio for poor outcome (failure to survive) in the group with the poor prognosis signature as compared with the good prognosis signature was 4.7 (95% confidence interval 2.8 – 7.9).

Our 249 gene signature overlapped by 11 genes with the 70 gene signature of van’t Veer et al. (16) (including NUSAP1, UCHL5, RAMP, DC13, PRC1, COL4A2, PITRM1, CENPA, MELK, KNTC2, and MCM6) and by 7 genes with the 76 gene
signature of Wang et al (33) (including MTB, POLQ, SUPT16H, FEN1, DUSP4, PLK1, and SMC4L1). The van’t Veer and Wang signatures overlapped by a single gene (cyclin E2). Our 19 gene signature had no genes in common with either of the previously published signatures.

Our 19-gene signature included several genes encoding proteins with roles in the cell cycle and in cell division. Cell-cycle genes were previously identified as important markers of prognosis in ER-positive younger patients (34). In this earlier study, 50 cell-cycle-related genes divided 83 ER-positive younger patients into two groups of good versus poor prognosis. The overall 10-year survival rates were 46% and 96% for the poor and good prognosis groups, respectively. Similarly, we found that a core group of predominantly cell cycle and mitotic organizing center genes (CDKN3, RRM3, FLJ10540, FOXM1, STK6, TRIP13, EIF4A1, FLJ10036, VRK1, TUBG1, CKS2, FLJ10517) made a strong contribution to stratifying tumors into good versus poor prognostic groups. One gene from our 19-gene signature, STK6, which encodes Aurora-A (35, 36) was also included in the 50-gene signature of Dai et al. (34), while 15 genes from our 249-gene signature were included in the Dai et al. signature (including BM039, DFKZp762E1312, LOC51203, LOC51659, ID-GAP, KNSL6, PRC1, STK6, CDC45L, SNRPA1, H2AFZ, CENPA, CDC6, BIRC5, and BLM). In addition to cell-cycle genes, our prognostic genes also encoded products with other functions, including genes involved in cytoskeletal regulation (ACTB, ACTN1) (37), cell survival (TNFRSF6B) (38, 39), polyamine biosynthesis (ODC1) (40), and cell-cell interactions (EPHA2) (41). The genes in these additional functional groups were important in subdividing the patients into subgroups with differing survival rates.
In conclusion, we report that the gene expression changes that commonly occur in non-malignant HMEC grown in 3D lrECM cultures provide gene expression signatures that effectively stratify patients into prognostic groups according to overall survival rates. Our 249 gene signature achieved a hazard ratio of 4.7, which is comparable to hazard ratios achieved by large scale supervised breast cancer microarray studies. Our results underscore the relevance of 3D lrECM cultures for studies of malignant transformation, and suggest potentially valuable new biomarkers for further clinical evaluation.

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Tables

**Table 1. Number of potential prognostic markers for breast cancer among the genes that were differential in the 3D time course (Method 1).**

<table>
<thead>
<tr>
<th>Total no. of genes</th>
<th>Number of significantly* differential genes when survival</th>
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</thead>
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<tr>
<td></td>
<td>(5 yrs)</td>
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<tr>
<td>All genes</td>
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<tr>
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<td>Random list 4</td>
<td>20</td>
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<td>Random list 5</td>
<td>20</td>
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</tbody>
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*Student's t-test, p<0.05.

#Total number reflects genes included on the Rosetta microarrays; hence values in some cases are less than the total number included on the Affymetrix microarrays.

Bold: > 50%.
FIGURE LEGENDS

Figure 1. HMEC cultured in lrECM form polarized structures and arrest growth in G0/G1. A. Phase-contrast image of typical acinar structures formed by 184 cells at day 7 in lrECM. Structures reach dimensions of 20-40 µm diameter. Bars equal 25 µm. B. Indirect immunofluorescent image showing basal polarity of alpha-6 integrin (green) in 184 cells at day 7 in lrECM. Cell nuclei were stained with DAPI (red). C. Flow cytometric analyses of propidium iodide stained S1 cells indicated that the majority of the cells accumulated in the G0/G1 phase of the cell cycle by day 7 in lrECM. D. Western blot analyses of cell-cycle regulatory molecules in S1 cells at days 5, 7, 10, and 15 after suspension in lrECM indicated that total as well as specific phosphorylated forms (serines 807/811, serine795, serine 612, threonine 821) of Rb decreased, while p27 increased, over the time course. Ponceau staining indicated that equivalent amounts of total protein were loaded in each lane.

Figure 2. A scheme is provided outlining Method 1 used to select sets of temporally co-regulated genes.

Figure 3. Sixty genes were determined by Method 1 to show significant changes in expression in both HMEC specimens during the time course of culture in lrECM. A. Temporal pattern of growth arrest in S1 and HMEC184 cells. The percentage of cells in S-phase decreased significantly between days 3 and 7. B. Expression levels of the 60 genes that were coordinately expressed in both cell types. The genes were grouped into categories based on whether they showed coordinate up-regulation early, up-regulation
late, down-regulation early, or down-regulation late during the time course (see text for details). Note that the scale is logarithmic. C. Changes in the expression levels of the 60 individual genes are plotted and organized by hierarchical cluster analysis. Red = up-regulated, blue = down-regulated genes. Complete gene names and GenBank IDs are available (see supplementary material).

Figure 4. Fourteen of the nineteen genes down-regulated late during acinar morphogenesis showed significant correlations with patient survival. 295 patients were grouped into quartiles based on the relative expression of each selected gene in their corresponding tumors. Survival of each quartile was plotted according to the method of Kaplan and Meier. P-values represent the outcomes of the log-rank tests between the upper and lower quartiles.

Figure 5. The set of 19 genes down-regulated late during acinar morphogenesis can be used to accurately cluster 295 breast cancer samples into prognostic groups. The rows represent the relative expression levels of the genes listed on the right. The columns represent individual patients. The scale of relative expression levels is the same as that shown in Figure 3C. Genes and tumor samples were arranged by a hierarchical cluster analysis using a Pearson metric. Dendrograms at the top and left reflect the degree of relatedness of the samples and genes, respectively. Dendrogram branch colors at the top indicate different prognostic groups. Genes in bold were significantly associated with survival (Kaplan-Meier analysis, p < 0.05, Figure 4). The panel below the cluster diagram indicates clinical parameters for the 295 patient samples. The black regions represent the
the percentages of patients surviving 10 years in each of the five prognostic groups are shown below.

Figure 6. A second set of 249 genes down-regulated late in the time course of HMECs in 3D lrECM cultures classifies good versus poor prognosis in breast cancer patients.  A. A scheme is provided outlining Method 2 used to select sets of temporally co-regulated genes. B. The set of 249 down-regulated genes identified by Method 2 clustered the 295 breast cancer samples into two prognostic groups. Genes are arranged by a hierarchical cluster analysis using a Pearson metric and samples are arranged using a distance metric. Clinical parameters are indicated below the diagram. C. Overall survival in the two groups was plotted by the method of Kaplan and Meier. P-value represents the outcome of the log-rank tests between the good and poor prognosis groups.
Figure 1. Fournier et al.
**Identification of temporally co-regulated genes (Method 1):**

<table>
<thead>
<tr>
<th></th>
<th>1.5-fold up# (2180 genes)</th>
<th>ANOVA, time (p&lt;0.05, error model)</th>
<th>Separate early vs late (161 vs 73 genes)</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1.5-fold down#@ (2652 genes)</td>
<td>ANOVA, time (p&lt;0.05, error model)</td>
<td>Separate early vs late (69 vs 282 genes)</td>
</tr>
<tr>
<td>184</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1.5-fold up# (1035 genes)</td>
<td>ANOVA, time (p&lt;0.05, error model)</td>
<td>Separate early vs late (327 vs 36 genes)</td>
</tr>
<tr>
<td></td>
<td>1.5-fold down#@ (1625 genes)</td>
<td>ANOVA, time (p&lt;0.05, error model)</td>
<td>Separate early vs late (76 vs 41 genes)</td>
</tr>
</tbody>
</table>

Identify temporally co-regulated gene classes:

- **UP EARLY in both cells** (21 genes)
- **UP LATE in both cells** (11 genes)
- **DOWN EARLY in both cells** (6 genes)
- **DOWN LATE in both cells** (22 genes)

---

# Genes at least 1.5-fold differential in at least 1 of 6 samples (2 experimental duplicates per time point).
@ The GeneSpring Cross Gene Error Model works by fitting a smooth curve to replicate variability resulting in error estimates that are based on pooling variability information for genes at similar expression levels. Since this method preferentially identifies positively changing (up-regulated) genes, the mean of the 7 day samples was used as the reference in cases where we wished to identify down-regulated genes. In all other cases, the mean of the 3 day samples was used as the reference.

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Figure 2. Fournier et al.
Figure 3. Fournier et al.
Figure 4. Fournier et al.
Figure 5. Fournier et al.
A  Identification of temporally co-regulated genes (Method 2):

- **S1**
  - 1.5-fold up* (2175 genes)
  - 1.5-fold down* (2356 genes)
  - 1.5-fold differential in either cell (5487 genes)

- **184**
  - 1.5-fold up* (1035 genes)
  - 1.5-fold down* (1459 genes)

Identify temporally co-regulated gene classes:
- UP EARLY in both cells (497 genes)
- UP LATE in both cells (79 genes)
- DOWN EARLY in both cells (142 genes)
- DOWN LATE in both cells (287 genes)

*Genes at least 1.5-fold differential in at least 1 of the later time points, 5 and 7 days.

B  Identification of temporally co-regulated genes (Method 2):

- **Good prognosis**
- **Poor prognosis**

C  Survival analysis:

- **Good**
- **Poor**

**Survival**

- 90% (138/154) survived 5 yrs
- 89% (83/141) survived 10 yrs

**P = 2.7e-10**

Figure 6. Fournier et al.