Featured Article

Gene Expression Profiles Predict Survival and Progression of Pleural Mesothelioma

Harvey I. Pass, Zhandong Liu, Anil Wali, Raphael Bueno, Susan Land, Daniel Lott, Fauzia Siddiq, Fulvio Lonardo, Michele Carbone, and Sorin Draghici

1Thoracic Oncology, Karmanos Cancer Institute, 2Department of Computer Science, and 3Center for Molecular Medicine and Genetics, Wayne State University, Detroit, Michigan; 4Department of Pathology, Harper University Hospital, Detroit, Michigan; 5Department of Pathology, Cardinal Bernardin Cancer Center, Loyola University, Maywood, Illinois; and 6Division of Thoracic Surgery, Brigham and Women’s Hospital, Harvard Medical School, Boston, Massachusetts

Abstract

Purpose: Clinical outcomes for malignant pleural mesothelioma (MPM) patients having surgery are imprecisely predicted by histopathology and intraoperative staging. We hypothesized that gene expression profiles could predict time to progression and survival in surgically cytoreduced pleural mesothelioma of all stages.

Experimental Design: Gene expression analyses from 21 MPM patients having cytoreductions and identical postoperative adjuvant therapy were performed using the U95 Affymetrix gene chip. Using both dChip and SAM, neural networks constructed a common 27 gene classifier, which was associated with either the high-risk and low-risk group of patients. Data were validated using real-time PCR and immunohistochemical staining. The 27 gene classifier was also used for validation in a separate set of 17 MPM patients from another institution.

Results: The groups predicted by the gene classifier recapitulated the actual time to progression and survival of the test set with 95.2% accuracy using 10-fold cross-validation. Clinical outcomes were independent of histology, and heterogeneity of progression and survival in early stage patients was defined by the classifier. The gene classifier had a 76% accuracy in the separate validation set of MPMs.

Conclusions: These data suggest that pretherapy gene expression analysis of mesothelioma biopsies may predict which patients may benefit from a surgical approach.

Introduction

Malignant pleural mesothelioma (MPM) is an orphan disease of which the management has defined curative options (1). The incidence of MPM worldwide is increasing, and the estimated number of deaths over the next 30 years from MPM in Western Europe alone could approach 250,000 (2). Our understanding of the pathophysiology of the disease, however, has improved with the advent of molecular investigations of the downstream effects of asbestos (3), as well as the realization that SV40 is linked with the disease in >60% of the cases (4). Studies of the molecular genetics of the disease have revealed the importance of signal transduction pathways including the epidermal growth factor receptor (5) and activator protein (6), as well as asbestosis-SV40 interactions, which could explain such phenomena as increased angiogenesis (7), telomerase (8, 9), MET overexpression (10, 11), abrogation of p53-induced apoptosis (12), and abnormalities of immune surveillance (13) among other findings.

There have been few global investigations using gene array technologies of the multiple genetic events in MPM due to the lack of uniformly treated patients with carefully documented survival and recurrence data, and the absence of large archives of banked frozen tissue with adequately preserved RNA. In addition to the gene-based classification (14), the ability to prognosticate on the basis of gene expression data from biopsy harvests could have important therapeutic implications for this group of unfortunate patients. Our hypothesis is that gene arrays performed on operative specimens from uniformly treated mesothelioma patients could predict survival and time to progression, and could cluster groups of patients as good or poor risk. Our test set population included stages I-III MPM patients resected by one surgeon. We analyzed our gene expression data using two programs, significance analysis microarrays (SAM) and dCHIP, to see whether concordant genes from both analyses could correctly cluster the patients into good or poor risk cohorts.

Materials and Methods

Patient Population. Tumor and normal tissues from 21 patients with MPM treated at the National Cancer Institute and the Karmanos Cancer Institute between August 1993 and May 2001 were used as the test set for the studies described below. Consent was received and the project approved by the local Institutional Review Boards. All of the patients had cytoreductive surgery, mediastinal lymph node dissection with staging by the International Mesothelioma Interest Group staging system (15), followed by postoperative cisplatin-based chemotherapy. All of the patients were followed from the surgery until death or through February 2003 with computerized tomography of the chest every 3 months. Interval change, which suggested progression of the MPM, was documented whenever possible by histological confirmation of disease. Tumors were classified as having epithelial, sarcomatoid, or mixed histology MPM by two
Gene Expression Prognostication for MPM

Calumenin, and IGFBP5. cDNAs were synthesized from 3
confirm differential gene expression profile of Plasmolipin,
Real-time quantitative reverse transcription-PCR was used to
compiled. dChip and SAM were used for expression profiling.

Real-Time Quantitative Reverse Transcription-PCR.
Real-time quantitative reverse transcription-PCR was used to
confirm differential gene expression profile of Plasmolipin,
Calumenin, and IGFBP5. cDNAs were synthesized from 3 µg
total RNA using an oligodeoxynucleotide primer and the
Superscript II reverse transcription kit (Life Technologies, Inc.).
The primers used were: Plasmolipin sense, 5'ATGTTC-GTCGCTGTCTTTCC-3'; antisense, 5'-ATCATCACCACAAC-ACGCAA-3'; Calumenin sense, 5'-GGCTGGATAATTACGAG-GATGT-3'; antisense 5'-GCAGGAAAGCTGTGAACTCC-3';
IGFBP5 sense 5'-GAGCTGAAGGCTGAAGCAGT-3', anti-

The amplified PCR products were 248 bp Plasmolipin
fragment, a 235 bp Calumenin fragment, and a 237 bp IGFBP5
fragment. Reactions containing 1 µg cDNA, SYBR green se-
sequence detection reagent, and sense and antisense primers were
assayed on an ABI 7700 sequence detection system (Applied Biosystems).
The PCR conditions were one cycle at 50°C for 2
min, one cycle at 95°C for 10 min, and 40 cycles of 95°C for 15
s, 60°C for 1 min. The accumulation of PCR product was
measured in real time as the increase in SYBR green fluores-
tence, and the data were analyzed using the Sequence Detector
program v1.6.3 (Applied Biosystems). Standard curves relating
initial template copy number to fluorescence and amplification
cycle were generated using the amplified PCR product as a
template and were used to calculate gene concentration (nor-
malized to β-actin values) in nM/µg cDNA.

Immunohistochemical Staining. Dparaffinized sec-
tions were stained manually, using the avidin-biotin complex
technique. In brief, 4 µm sections were collected on gelatin-
coated slides, deparaffinized, and rehydrated via passage
through graded alcohol solutions, according to standard histo-
logical techniques. Heat-induced epitope retrieval was obtained
by heating slides for 20' in a citrate buffer (2 mm citric acid/10
mm Na citrate; pH 6) in a steam bath, followed by cooling down
for 20' at room temperature. Slides were presaturated with
0.05% BSA in PBS for 10', quenched in 2% H2O2 for 10', and
preincubated with 0.5% nonimmune goat serum (Sigma) for 10'
at room temperature. The sections were then incubated with the
primary antibodies. Antibodies used, dilutions, and incubation
conditions were monoclonal Nm 23 antibody (Santa Cruz Bio-
technology-H1; sc 465) 1:20 dilution, overnight incubation at
4°C; monoclonal Integrin α 6 (Santa Cruz Biotechnology BQ;
sc-13542) 1:10 dilution, overnight incubation at 4°C; and poly-
clonal FGF-7 (Santa Cruz Biotechnology-C19; sc-1365), 1:40
dilution, 2-h incubation at room temperature. After three washes
in BSA/PBS (5' each) localization of the primary antibody and
color development were obtained by incubation with biontyl-
lated antiamous (monoclonal antibodies) or antigen (FGF-7)
secondary antibodies (Vector), followed by incubation with streptavidin-coupled horseradish peroxidase, using an avidin-
biotin detection kit, with AEC as the chromogen (Vector Labs,
Burlingame, CA). Sections were lightly counterstained with
hematoxylin (Sigma) and coverslipped. Positive controls were
included with each run and included colon cancer for Nm23,
normal skin for Integrin α 6, and ductal carcinoma of breast for
FGF-7. Negative controls were included with each run by omit-
ting the primary antibody. Score was separately assessed for
intensity and extent of positivity. Intensity was scored on a scale
of 1 (lightest) to 3 (strongest). The extent of tumor cells posi-
tivity was scored as negative, no stain; focal, ≤10% positive

tumor cells; moderate, >10% positive cells ≤50%; and diffuse,
≥50% positive tumor cells. Positive stain occurred in the cyto-

Table 1 Mesothelioma test set characteristics

<table>
<thead>
<tr>
<th>Survivor group</th>
<th>Number</th>
<th>Sex</th>
<th>IMIG Stage</th>
<th>Histology</th>
<th>Survival (mos.)</th>
<th>Status</th>
<th>Time to recurrence</th>
<th>Progression</th>
</tr>
</thead>
<tbody>
<tr>
<td>Short-term</td>
<td>1</td>
<td>M</td>
<td>III</td>
<td>Epithelial</td>
<td>2</td>
<td>D</td>
<td>1.0</td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>M</td>
<td>II</td>
<td>Epithelial</td>
<td>3.6</td>
<td>D</td>
<td>1.8</td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>M</td>
<td>III</td>
<td>Epithelial</td>
<td>5.2</td>
<td>D</td>
<td>3.4</td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td>8</td>
<td>M</td>
<td>III</td>
<td>Epithelial</td>
<td>3.2</td>
<td>D</td>
<td>3.2</td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td>12</td>
<td>M</td>
<td>III</td>
<td>Biphasic</td>
<td>4.3</td>
<td>D</td>
<td>3.0</td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td>17</td>
<td>M</td>
<td>III</td>
<td>Epithelial</td>
<td>4.8</td>
<td>D</td>
<td>1.64</td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td>39</td>
<td>M</td>
<td>II</td>
<td>Biphasic</td>
<td>2</td>
<td>D</td>
<td>1.5</td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td>38</td>
<td>M</td>
<td>III</td>
<td>Biphasic</td>
<td>2</td>
<td>D</td>
<td>1.75</td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td>30</td>
<td>F</td>
<td>III</td>
<td>Epithelial</td>
<td>7.3</td>
<td>D</td>
<td>2.2</td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td>29</td>
<td>M</td>
<td>III</td>
<td>Biphasic</td>
<td>1.7</td>
<td>D</td>
<td>1.3</td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td>33</td>
<td>M</td>
<td>II</td>
<td>Epithelial</td>
<td>10</td>
<td>D</td>
<td>8</td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>M</td>
<td>III</td>
<td>Epithelial</td>
<td>48</td>
<td>A</td>
<td>17</td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>M</td>
<td>I</td>
<td>Epithelial</td>
<td>95</td>
<td>A</td>
<td>95</td>
<td>N</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>F</td>
<td>III</td>
<td>Biphasic</td>
<td>25</td>
<td>D</td>
<td>22</td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td>7</td>
<td>M</td>
<td>III</td>
<td>Biphasic</td>
<td>32</td>
<td>D</td>
<td>14</td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td>11</td>
<td>M</td>
<td>I</td>
<td>Epithelial</td>
<td>95</td>
<td>A</td>
<td>14</td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td>16</td>
<td>M</td>
<td>II</td>
<td>Epithelial</td>
<td>99</td>
<td>A</td>
<td>15</td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td>19</td>
<td>F</td>
<td>III</td>
<td>Sarcomatoid</td>
<td>103</td>
<td>A</td>
<td>6</td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td>21</td>
<td>F</td>
<td>I</td>
<td>Epithelial</td>
<td>102</td>
<td>A</td>
<td>60</td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td>22</td>
<td>F</td>
<td>III</td>
<td>Epithelial</td>
<td>28</td>
<td>D</td>
<td>12</td>
<td>Y</td>
</tr>
<tr>
<td></td>
<td>15</td>
<td>M</td>
<td>II</td>
<td>Epithelial</td>
<td>58</td>
<td>D</td>
<td>12</td>
<td>Y</td>
</tr>
</tbody>
</table>

Downloaded from clincancerres.aacrjournals.org on January 27, 2018. © 2004 American Association for Cancer Research.
Data Analysis and Neural Network Methods. *T* tests were used to identify differences in mean gene expression levels between comparison groups. We constructed of the classifiers using a multilayer feedforward backpropagation neural network. A three layer feed forward artificial neural network was trained based on the expression value of differentially regulated genes, and the accuracy of the neural network was confirmed with 10-fold cross-validation. The 10-fold cross-validation involves constructing 10 different classifiers in 10 training runs. Each training run divides the available data into a training set, including 90% of the data and a validation set including 10% of the data. The classifier is constructed based on the training set and tested on the validation set. Each training run leaves out different patterns such as at the end of the entire process, each

![Fig. 1](image)

A, gene expression patterns determined using hierarchical clustering of the 21 mesothelioma patients against the top 95 genes identified by dChip. Substantially elevated (red) or decreased (blue) gene expression of the genes is observed in individual tumors. *B*, gene expression patterns determined using hierarchical clustering of the 21 mesothelioma patients against the 27 common genes between dChip and SAM. Two groups are defined, which have contrasting gene expression patterns with the exception of 3 genes, which are overexpressed in the majority of cases (see “Discussion”).
available patient will have been left out at least once. The performance reported represents an average of these 10 training runs. The process ensures that the resulting classifiers are always tested on data that has never been used during the training, thus testing its generalization abilities. We calculated the intersection of the genes selected by the dChip and Sam, and constructed a similar neural network classifier based on only those genes that were selected by both methods. To inspect visually the expression profiles of the genes, a hierarchical clustering was constructed for each group of genes using average linkage and standardized Euclidean distance.

The performance of each classifier was assessed in two ways. Firstly, each classifier was tested and assessed using 10-fold cross-validation, as described above. Secondly, we used the trained classifier on a previously unseen data set of 17 mesothelioma patients including 8 long-term survivors and 9 short-term survivors. These data were obtained from the Division of Thoracic Surgery of the Brigham and Women’s Hospital (16). Kaplan Meier survival plots and log rank tests were used to assess differences between poor and good risk groups using MedCalc Software (Mariakerke, Belgium).

Onto-Express (OE) Analysis. Using OE (17), we performed a functional analysis to identify the main biological processes and pathways involving these prognostic genes. OE is a tool designed to mine the available functional annotation data and find relevant biological processes. The input to OE is the list of GenBank accession numbers, Affymetrix probe IDs, or UniGene cluster IDs. OE constructs a functional profile for each of the Gene Ontology categories (18), including cellular component, biological process, and molecular function, as well as biochemical function and cellular role, as defined by Proteome. As biological processes can be regulated within a local chromosomal region (e.g., imprinting), an additional profile is constructed for the chromosome location. Statistical significance values are calculated for each category using either a hypergeometric or binomial distribution depending on the number of genes on the array used.

Results

Hierarchical Profile Clustering Yields Two Mesothelioma Subsets. Using the U95 Affymetrix oligonucleotide arrays, we generated profiles for 21 MPMs, which had been defined as “short-term” and “long-term” groups based on their actual survival from operation as either < or >12 months (Table 1). Median survival time for the short-term group was 3.6 months (range, 2–10 months) compared with 58 months (range, 25–102 months) for the long-term group ($P < 0.0001$). The time from surgery to radiographic progression of disease was a median of 1.8 months (range, 1–8 months) and 14.5 months (range, 6–95 months) for the short- and long-term groups, respectively ($P < 0.0001$).

Selection of Differentially Regulated Genes: dChip. The dChip software package (19) normalized the 21 Affymetrix chips by the chip with median expression value. The unsupervised clustering on all 12,625 of the genes produced a dendrogram in which the gene expression profiles of the long- and short-term survivors were not consistently different. A PM/MM model-based normalization was performed to select the genes that were differentially regulated between the long- and short-term groups. The performance of each classifier was assessed in two ways. Firstly, each classifier was tested and assessed using 10-fold cross-validation, as described above. Secondly, we used the trained classifier on a previously unseen data set of 17 mesothelioma patients including 8 long-term survivors and 9 short-term survivors. These data were obtained from the Division of Thoracic Surgery of the Brigham and Women’s Hospital (16). Kaplan Meier survival plots and log rank tests were used to assess differences between poor and good risk groups using MedCalc Software (Mariakerke, Belgium).

Onto-Express (OE) Analysis. Using OE (17), we performed a functional analysis to identify the main biological processes and pathways involving these prognostic genes. OE is a tool designed to mine the available functional annotation data and find relevant biological processes. The input to OE is the list of GenBank accession numbers, Affymetrix probe IDs, or UniGene cluster IDs. OE constructs a functional profile for each of the Gene Ontology categories (18), including cellular component, biological process, and molecular function, as well as biochemical function and cellular role, as defined by Proteome. As biological processes can be regulated within a local chromosomal region (e.g., imprinting), an additional profile is constructed for the chromosome location. Statistical significance values are calculated for each category using either a hypergeometric or binomial distribution depending on the number of genes on the array used.

Results

Hierarchical Profile Clustering Yields Two Mesothelioma Subsets. Using the U95 Affymetrix oligonucleotide arrays, we generated profiles for 21 MPMs, which had been defined as “short-term” and “long-term” groups based on their actual survival from operation as either < or >12 months (Table 1). Median survival time for the short-term group was 3.6 months (range, 2–10 months) compared with 58 months (range, 25–102 months) for the long-term group ($P < 0.0001$). The time from surgery to radiographic progression of disease was a median of 1.8 months (range, 1–8 months) and 14.5 months (range, 6–95 months) for the short- and long-term groups, respectively ($P < 0.0001$).

Selection of Differentially Regulated Genes: dChip. The dChip software package (19) normalized the 21 Affymetrix chips by the chip with median expression value. The unsupervised clustering on all 12,625 of the genes produced a dendrogram in which the gene expression profiles of the long- and short-term survivors were not consistently different. A PM/MM model-based normalization was performed to select the genes that were differentially regulated between the long- and short-term groups. The performance of each classifier was assessed in two ways. Firstly, each classifier was tested and assessed using 10-fold cross-validation, as described above. Secondly, we used the trained classifier on a previously unseen data set of 17 mesothelioma patients including 8 long-term survivors and 9 short-term survivors. These data were obtained from the Division of Thoracic Surgery of the Brigham and Women’s Hospital (16). Kaplan Meier survival plots and log rank tests were used to assess differences between poor and good risk groups using MedCalc Software (Mariakerke, Belgium).

Onto-Express (OE) Analysis. Using OE (17), we performed a functional analysis to identify the main biological processes and pathways involving these prognostic genes. OE is a tool designed to mine the available functional annotation data and find relevant biological processes. The input to OE is the list of GenBank accession numbers, Affymetrix probe IDs, or UniGene cluster IDs. OE constructs a functional profile for each of the Gene Ontology categories (18), including cellular component, biological process, and molecular function, as well as biochemical function and cellular role, as defined by Proteome. As biological processes can be regulated within a local chromosomal region (e.g., imprinting), an additional profile is constructed for the chromosome location. Statistical significance values are calculated for each category using either a hypergeometric or binomial distribution depending on the number of genes on the array used.

Results

Hierarchical Profile Clustering Yields Two Mesothelioma Subsets. Using the U95 Affymetrix oligonucleotide arrays, we generated profiles for 21 MPMs, which had been defined as “short-term” and “long-term” groups based on their actual survival from operation as either < or >12 months (Table 1). Median survival time for the short-term group was 3.6 months (range, 2–10 months) compared with 58 months (range, 25–102 months) for the long-term group ($P < 0.0001$). The time from surgery to radiographic progression of disease was a median of 1.8 months (range, 1–8 months) and 14.5 months (range, 6–95 months) for the short- and long-term groups, respectively ($P < 0.0001$).

Selection of Differentially Regulated Genes: dChip. The dChip software package (19) normalized the 21 Affymetrix chips by the chip with median expression value. The unsupervised clustering on all 12,625 of the genes produced a dendrogram in which the gene expression profiles of the long- and short-term survivors were not consistently different. A PM/MM model-based normalization was performed to select the genes that were differentially regulated between the long- and short-
term survivor groups with the following criteria: (a) E-B > 100; (b) E/B > 1.2 or B/E > 1.2; and (c) two group t test has a P < 0.05. E and B stand for the average expression value in the experiment (long-term survivors, good risk) and control (short-term survivors, poor risk) groups, respectively. Ninety-five genes satisfied the two criteria above (Fig. 1A). These 95 genes were used to construct an artificial neural network with a 95-14-2 architecture. This network achieved an accuracy of 90.5% measured by 10-fold cross-validation. The sensitivity and specificity of this artificial neural network classifier were 90% and 90.9%, respectively.

Selection of Differentially Regulated Genes: SAM. We analyzed the same data using the Significance Analysis of Microarray (20) using a fold change of 1.2 and a delta value of 0.6679. Because of the sensitivity of setting parameters in SAM, we could only select 100 genes (as opposed to 95 in dChip) with a false discovery rate of 26.6%. A neural network constructed a classifier based on these SAM-selected genes with an accuracy of 100% (achieved on 10-fold cross-validation). The sensitivity and specificity of this artificial neural network classifier were 100% and 100%, respectively.

Common Genes between dChip and SAM. To define which important genes were common to dChip and SAM, an unpaired two group comparison was performed on the dChip normalized data. Interestingly, the intersection of the 95 genes selected by dChip with the 100 genes selected by SAM yielded only 27 common genes. This suggested that: (a) the data were rather noisy and the subset of good predictors might be <95 genes, and (b) at least one but possibly both methods picked up a number of genes that were less than ideal predictors. To investigate these hypotheses, we built a classifier based on the genes common between the two sets (i.e., 95 dChip genes and...
100 SAM genes as opposed to the most significant 27 in each analysis). The clustering dendrogram for these common 27 genes is seen in Fig. 1B. This classifier yielded an accuracy of 95.2% on the validation dataset with 10-fold cross-validation. The confusion matrix is shown in Table 2.

**Real-Time PCR and Immunohistochemistry Analyses.** Six genes of the 27 were arbitrarily selected to verify the microarray expression data. Real-time quantitative reverse transcription-PCR was performed using primers for Plasmolin, Calumenin, and IGFBPS on a minimum of 7 tumor samples. As seen in Fig. 2, there was excellent correlation between the array expression data and the real-time PCR, with the best correlation seen for IGFBPS (0.99); those for plasmolin and calumenin were 0.86. Immunohistochemical analysis of integrin α6, fibroblast growth factor, and IGFBPS was performed for patients 4 and 39, each representing the two clusters to determine whether RNA expression was reflected appropriately by the corresponding proteins in tumors (Fig. 3). When one compares the expression of the proteins by immunohistochemistry with the clustergram (Fig. 1B), it is seen that patient 4 with a more favorable prognosis had elevated levels of integrin and FGF7, and both of these proteins were scored as having a stronger and more focal intensity with more cells staining in patient 4 than patient 39. Staining for NM23A revealed approximately equal intensity and quantity of cellular staining.

**Biological Processes Impacted by MPM.** We used OE (21) to identify the biological processes impacted significantly by the 27 differentially regulated genes found by both analysis approaches (Table 3). These biological processes include cell proliferation (P = 0.016), lipid metabolism (P = 0.001), positive regulation of cell proliferation (P = 0.025), cell-cell signaling (P = 0.095), regulation of transcription from Pol II promoter (P = 0.071), pathogenesis (P = 0.010), cell cycle arrest (P = 0.003), negative regulation of cell proliferation (P =

<table>
<thead>
<tr>
<th>Table 3</th>
<th>The 27 common dChip and SAM genes for malignant pleural mesothelioma prognostication</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gene name</td>
<td>Chromosomal location</td>
</tr>
<tr>
<td>Clusterin (complement lysis inhibitor, SP-40, apolipoprotein J)</td>
<td>8p21-p12</td>
</tr>
<tr>
<td>Transmembrane 4 superfamily member 11 (plasmolin)</td>
<td>1q13</td>
</tr>
<tr>
<td>Cysteine-rich protein 1 (intestinal)</td>
<td>7q11.23</td>
</tr>
<tr>
<td>Dishevelled associated activator of morphogenesis 1</td>
<td>14q22.3</td>
</tr>
<tr>
<td>Homo sapiens mRNA; cDNA DKFZp564B076 (from clone DKFZp564B076)</td>
<td>Ch. 6</td>
</tr>
<tr>
<td>Alcohol dehydrogenase IB (class I), β polypeptide</td>
<td>4q21-q23</td>
</tr>
<tr>
<td>Human insulin-like growth factor binding protein 5 (IGFBPS)</td>
<td>Ch. 2</td>
</tr>
<tr>
<td>DKFZPS86A0522 protein</td>
<td>1q21</td>
</tr>
<tr>
<td>CD27-binding (Siva) protein</td>
<td>1q21.3</td>
</tr>
<tr>
<td>Lutheran blood group (Auerberb antigen included)</td>
<td>19q13.2</td>
</tr>
<tr>
<td>Short-chain dehydrogenase/reductase 1</td>
<td>1p36.1</td>
</tr>
<tr>
<td>Heterogeneous nuclear ribonucleoprotein H1 (H)</td>
<td>Ch. 3</td>
</tr>
<tr>
<td>Homo sapiens, similar to Y43E12A2.p, clone MGC:33537</td>
<td>Ch. 10</td>
</tr>
<tr>
<td>IMAGE:4821347</td>
<td>35369_at</td>
</tr>
<tr>
<td>KIAA0937 protein</td>
<td>1q21.3</td>
</tr>
<tr>
<td>Monoglyceroide lipase</td>
<td>10pter-q25.3</td>
</tr>
<tr>
<td>KIAA0275 gene product</td>
<td>1q21</td>
</tr>
<tr>
<td>BTG family, member 2</td>
<td>7q22</td>
</tr>
<tr>
<td>Calumenin</td>
<td>7q22</td>
</tr>
<tr>
<td>Selenium binding protein 1</td>
<td>2q21-q22</td>
</tr>
<tr>
<td>Acidic (leucine-rich) nuclear phosphoprotein 32 family, member B</td>
<td>2q21</td>
</tr>
<tr>
<td>KIAA1237 protein</td>
<td>3q21.3</td>
</tr>
<tr>
<td>Integrin, α6</td>
<td>10q23.32</td>
</tr>
<tr>
<td>Adipocyte specific 2</td>
<td>2p24-q22</td>
</tr>
<tr>
<td>V-myc myelocytomatosis viral oncogene homolog (avian)</td>
<td>8q24.12-q24.13</td>
</tr>
<tr>
<td>Nonmetastatic cells 1, protein (NM23A) expressed in</td>
<td>17q21.3</td>
</tr>
<tr>
<td>Fibroblast growth factor 7 (keratinocyte growth factor)</td>
<td>17q15-q21.1</td>
</tr>
</tbody>
</table>
0.034), inflammatory responses ($P = 0.022$), apoptosis induction by extracellular signals ($P = 0.001$), DNA damage response, activation of p53 ($P = 0.001$), and RNA processing ($P = 0.006$). The $P$ was calculated using a binomial distribution (21).

**Survival and Recurrence Prognostication: Test Set.**

The dendogram defined by the 27 common genes identified a "poor risk" and "good risk" group of MPMs within this test set ($P = 0.0018$). The median survival of the actual short-term survival group was 3.6 months and that of the predicted poor-risk group was 4.3 months (Fig. 4, A and B). No observed propensity for segregation of epithelial mesotheliomas into the good-risk group or other histologies into the poor-risk group was observed. Time-to-progression curves for the gene set also mirrored the test set actual clinical data (Fig. 4, C and D).

There were statistical differences in survival ($P = 0.0484$) between actual stage I (median survival not reached), stage II (median survival 10 months), and stage III (median survival 5.2 months) patients (Fig. 5, A and B). Prognostic heterogeneity, however, was detected within stages when survivals and progression were examined based on the gene classifier. When surgically staged I and II patients were grouped together, their survival was significantly different depending on which cluster the 27 gene classifier assigned them to. As seen in Fig. 5C, the patients in either stage I or 2 who were assigned to the poor-risk cluster were found to have significantly shorter survival than those assigned to the other cluster ($P = 0.0042$), and the time to progression of the stage I and II patients (Fig. 5D) was 1.8 months for those assigned to the poor-risk cluster and 15 months for those in the good-risk cluster ($P = 0.0042$). These differences were only suggestive for the stage III mesothelioma patients with regard to survival (Fig. 5, E and F). There was, however, a statistically longer time to progression for those stage III mesotheliomas who were clustered as good-risk (17 month median) compared with the poor-risk cluster (2.6 month median; $P = 0.0397$).

**Validation with an Independent Set of Pleural Mesotheliomas.** We assessed the performance of the 27 gene neural network classifier using oligonucleotide gene expression data obtained from a completely independent [Brigham and Women’s (B&W) Hospital] sample of 17 resected MPMs, all stages I or II. First, we trained our neural network with the genes selected by dChip from our dataset, and then the neural network was used to classify patients from B&W dataset. The classification accuracy is only 52.94%. The neural network built with genes selected by SAM from our dataset also did not yield a good result with a classification accuracy of only 47.05%. The classifier built with the common genes between dChip genes and SAM genes yielded a classification accuracy of 76.47% (Table 2), and based on our 27 important
genes, we applied hierarchical clustering on the patients from B&W dataset. The resulting dendogram revealed two groups (Fig. 6A) from the B&W data, and the predicted survival of the B&W patients (Fig. 6C) was similar to their actual (Fig. 6B) survival (median survival, 5.5 months versus 5 months) for the poor-risk groups. Moreover, the survival differences between the gene-predicted 5 good-risk patients in the B&W and the 12 poor-risk patients approached significance ($P = 0.0511$).

Discussion

Of the 27 genes found to be important in this investigation, 18 have been thoroughly classified in the literature, and few have been associated with MPM. Clusterin SP-40, acidic protein rich in leucine, and cysteine-rich protein were overexpressed in the vast majority of the MPMs, and, hypothetically, could be part of some pathway common to mesothelial carcinogenesis. Selenium binding protein was consistently overexpressed in good-risk patients, and was the only gene common to both the test set and validation set. SIVA or CD27-binding protein is part of an apoptotic pathway induced by CD27 antigen, a member of the tumor necrosis factor receptor superfamily. CD27 regulates the death and differentiation of T and B cells, and provides signals needed for the correct activation of specific T cells. Whether SIVA is important in MPMs for immune surveillance, and as such, could contribute to a longer survival, is unknown.

Short chain dehydrogenase/reductase (retSDR1) reduces all-trans-retinal during bleached visual pigment regeneration (22). In the absence of retSDR1, and under low concentrations of circulating retinol, the lack of production of vitamin A active...

![Gene expression in mesothelioma detects heterogeneity of natural history between patients with similar surgical stages.](image)
metabolites could contribute to cancer development and progression. Retinoic acid decreases the synthesis of fibronectin and laminin, and migration of MPM implying that retinoids and their binding may decrease MPM local invasion and tumor progression (23). 

\[ \text{BTG 2} \]

is a member of a group of structurally related antiproliferative proteins, which mediate a common signal transduction growth arrest and differentiation pathway. This pathway interacts with \( N \)-methyl transferase, the chief enzyme for post-translational modifications of proteins by protein methylation (24). BTG influences substrates of this methylation pathway including heterogeneous nuclear ribonucleoproteins (hnRNP), which was generally lower in expression in the good-risk group.

\[ \text{hnRNP H1} \]

influences pre-mRNA processing, and its role in cancer is unclear; however, hnRNP H1 has been identified as a protein that binds to the negative regulator spicing element of the Rous Sarcoma Virus (25, 26), and also binds a G-rich element downstream of the core SV40 late polyadenylation signal and stimulates 3' processing (27). These data suggest that hnRNP H1 could modulate late viral element activity in two known carcinogenic viruses, of which one (SV40) has now been implicated in the pathogenesis of MPM (4).

Calumenin, a calcium binding protein localized in the endoplasmic reticulum involved in protein folding and sorting, was uniformly underexpressed in the good-risk group. Calumenin has not specifically been associated with MPM before, but serum amyloid A and P have been noted to be connected with MPM (29–31). Nm23a, variably expressed in the good-risk group, was uniformly overexpressed in the poor-risk group. This transcription factor associated with highly metastatic cells (32) plays a role in myc expression. It has been shown that myc can up-regulate nm23-H1 and nm23-H2 expression (33), and data are variable regarding the prognostic significance of nm23 in human neoplasms (34–39).

Our data analysis is novel compared with other reports in the literature. Differences in array platforms, data analysis packages, or modifications in the filtering of the genes may result in differences in classifier genes among reports examining the same disease. We have demonstrated this phenomenon in our own work by comparing the top significant genes of interest from dChip and SAM, and found that there was not uniform agreement between the two programs on these genes, and a 10-fold cross-validation testing yielded differing results for the two programs. We feel that by selecting only those genes that were common to the two analyses, the resulting classifiers are more robust. Moreover, neither the top 27 genes from dChip or SAM could segregate survival differences among the B&W MPMs; however, the common set was 76% accurate in this independent set of specimens. It is likely that the patient population from the National Cancer Institute/Karmanos experience is different from that of the Boston experience. Our group

![Fig. 6](image_url)

**Validation of gene classifier in other surgically resected mesotheliomas.**

A, gene expression patterns determined using hierarchical clustering of the 17 mesothelioma patients against the 27 common genes between dChip and SAM. Two groups were defined, which have contrasting gene expression patterns similar to that seen in the test set (Fig. 1B). B, the actual survival time between the short and long term surviving patients in the B&W group was significant (\( P = 0.0012 \)). C, the 27 gene classifier defined two groups of mesotheliomas of which the difference in survival approached significance (\( P = 0.0511 \)).
included patients who were in International Mesothelioma Interest Group stage III, usually on the basis of metastases to the mediastinal lymph nodes at the time of surgery. The Boston MPMs were all stage I or II by the B&W classification, and in this classification, stage II patients may have intraparenchymal lymph node involvement, which would be stage III in the International Mesothelioma Interest Group classification. Moreover, the Boston MPM expression array data would be based on tumors that at the time of surgery did not have disease in the mediastinal lymph nodes (B&W classification stage III). Criteria selection for gene expression differed from the validation set in that the B&W group stipulated minimal expression of 500 and at least 2-fold differences. Given the huge differences in clinical presentation of these subsets, it is actually gratifying that a 76% validation of our gene set could be achieved in the Boston patients.

We have described a set of 27 genes that segregates good-risk and poor-risk surgically treated MPM patients with validation in an independent set of MPM patients. The classifier-predicted good-risk group had significantly longer survival and significantly longer time to progression than the poor-risk group. Moreover, these classifications were independent of the “conventional wisdom” that epithelial mesotheliomas have a better prognosis than biphasic and sarcomatoid histotypes. The data also reflect staging heterogeneity in relation to stage-related prognosis. Good-risk MPMs who were pathological stage I and II had strikingly different survival time and tendency to progression compared with stage I and II patients assigned to the poor-risk cohort. These data reflect the clinical scenario that certain patients with favorable prognostic demographics clinically (i.e., epithelial histology and early stage) nevertheless may recur quickly after operation despite maximal cytoreduction. Such data, after validation in larger numbers of patients, could be important to guide clinical decisions regarding appropriate therapy before committing the MPM patient to surgery.

Acknowledgments

The superb technical assistance of Valerie Murphy, Julia Sluchak-Carlsen, and Apurvi Desai was crucial for the successful completion of these series of studies.

References

Gene Expression Profiles Predict Survival and Progression of Pleural Mesothelioma

Harvey I. Pass, Zhandong Liu, Anil Wali, et al.


Updated version
Access the most recent version of this article at:
http://clincancerres.aacrjournals.org/content/10/3/849

Cited articles
This article cites 39 articles, 8 of which you can access for free at:
http://clincancerres.aacrjournals.org/content/10/3/849.full#ref-list-1

Citing articles
This article has been cited by 11 HighWire-hosted articles. Access the articles at:
http://clincancerres.aacrjournals.org/content/10/3/849.full#related-urls

E-mail alerts
Sign up to receive free email-alerts related to this article or journal.

Reprints and Subscriptions
To order reprints of this article or to subscribe to the journal, contact the AACR Publications Department at pubs@aacr.org.

Permissions
To request permission to re-use all or part of this article, use this link
http://clincancerres.aacrjournals.org/content/10/3/849.
Click on "Request Permissions" which will take you to the Copyright Clearance Center's (CCC) Rightslink site.