The molecular classification of multiple myeloma

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To better define the molecular basis of multiple myeloma (MM), we performed unsupervised hierarchic clustering of mRNA expression profiles in CD138-enriched plasma cells from 414 newly diagnosed patients who went on to receive high-dose therapy and tandem stem cell transplants. Seven disease subtypes were validated that were strongly influenced by known genetic lesions, such as c-MAF– and MAFB-, CCND1- and CCND3-, and MMSET-activating translocations and hyperdiploidy. Indicative of the deregulation of common pathways by gene orthologs, common gene signatures were observed in cases with c-MAF- and MAFB activation and CCND1 and CCND3 activation, the latter consisting of 2 subgroups, one characterized by expression of the early B-cell markers CD20 and PAX5. A low incidence of focal bone disease distinguished one and increased expression of proliferation-associated genes of another novel subgroup. Comprising varying fractions of each of the other 6 subgroups, the proliferation subgroup dominated at relapse, suggesting that this signature is linked to disease progression. Proliferation and MMSET-spike groups were characterized by significant overexpression of genes mapping to chromosome 1q, and both exhibited a poor prognosis relative to the other groups. A subset of cases with a predominating myeloid gene expression signature, excluded from the profiling analyses, had more favorable baseline characteristics and superior prognosis to those lacking this signature.

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Introduction

Multiple myeloma (MM) is a malignancy of antibody-secreting, terminally differentiated B cells that home to and expand in the bone marrow, with symptoms related to anemia, immunosuppression, bone destruction, and renal failure.1,2 Bone lesions developing adjacent to plasma cell foci result from the activation of osteoclasts and inactivation of osteoblasts.3-5 Many of the pathogenetic mechanisms of this clinically heterogeneous malignancy have been unraveled by application of molecular genetics.6-8 While sharing most of the genetic lesions seen in MM,9,11 monoclonal gammopathy of undetermined significance (MGUS) rarely progresses to overt MM.12 The universal activation of I of the 3 cyclin D genes is consistent with this being an initiating event in MM.13 Nonhyperdiploid MM, present in 40%, is characterized by transcriptional activation of CCND1, CCND3, MAF, MAFB, or FGFR3/MMSET genes (resulting from translocations involving the immunoglobulin heavy chain locus).8 While hyperdiploidy and CCND1 activation confer a favorable prognosis, MAF, MAFB, or FGFR3/MMSET activation and deletion of chromosomes 13 and 17 are associated with poor prognosis.14-25

Although high-dose therapy has markedly improved MM prognosis,26-28 individual patients’ survival remains variable29,30 and cannot be accurately predicted with current prognostic models.31,32 In lymphoma and leukemia, microarray profiling has helped establish clinically relevant disease subclassifications.33-41 In MM, such an approach has identified genes involved in pathogenesis and “drugable” pathways.3,21,42-50 Toward the development of a prognostically relevant molecular classification of MM, gene expression profiling was performed on CD138-enriched plasma cells from 414 newly diagnosed patients treated with high-dose melphalan-based tandem transplants.

Patients, materials, and methods

Patients: training and test sets

Gene expression profiling of highly purified bone marrow plasma cells was performed in 414 newly diagnosed patients with MM. The training set consisted of 256 cases enrolled on total therapy 2 (TT2),28 representing a subset of all 668 patients. There were similar baseline and outcome features in those with and without array profiling performed (data not shown). No difference was observed in event-free or overall survival among the 256 patients relative to the total population of 668, whether or not they were randomized to thalidomide (data not shown). With a median follow-up of 36 months, 110 disease-related events and 72 disease-related deaths have occurred. The test set comprised 158 patients enrolled in total therapy 3 (TT3) and served to validate the gene expression model developed with total therapy 2 samples; the short median follow-up of 12 months precluded an analysis of survival differences based on gene expression classes in the test set. Baseline clinical characteristics of training and test sets are

The online version of this article contains a data supplement.

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presented in Table S1, which is available on the Blood website (see the Supplemental Tables link at the top of the online article). The institutional review board of the University of Arkansas for Medical Sciences had approved the research studies, and all subjects had provided written informed consent to both treatment protocols and sample procurement, in accordance with the Declaration of Helsinki.

**Plasma cell selection and filtering of samples for analysis**

Plasma cells were enriched by anti-CD138 immunomagnetic bead selection of mononuclear cell fractions of bone marrow aspirates in a central laboratory. In all, 351 and 214 consecutive patients were evaluated in the training and test sets, respectively. All samples applied to microarray contained more than 85% plasma cells as determined by 2-color flow cytometry (CD38^dim^ and CD45^low^) performed after selection. We have previously reported that CD138 selection can result in cell populations with varying degrees of contamination with cells of the myeloid lineage and/or normal plasma cells. 13 To maintain consistency and ensure faithful assessment of the MM transcriptome, we eliminated samples with high degree of contamination of either of these 2 cell types as assessed by gene expression signatures. In all, 95 of the 351 training set samples and 56 of the 214 test set samples were excluded from the unsupervised clustering and subsequent gene expression–based classifications.

All primary microarray data presented in this paper have been deposited in the NIH Gene Expression Omnibus (GEO; National Center for Biotechnology Information [NCBI], http://www.ncbi.nlm.nih.gov/geo/) under accession number GSE2658.

**Gene expression profiling and data analyses**

Gene expression profiling was performed with the Affymetrix U133Plus2.0 microarray platform (Santa Clara, CA) using methods previously described.42 All data used in these analyses were derived with the Affymetrix Microarray Suite GCOS1.1 software. Affymetrix signals were transformed by the log-base 2 for each sample. The genes used in the analysis were chosen as follows. Genes having a present detection call in fewer than 3% of the samples and duplicate genes with a smaller standard deviation were selected using significance analysis of microarray (SAM)52 and chi-square correlation metric51 was used to identify disease subgroups.

Genes that were uniquely overexpressed or underexpressed in specific subgroups defined by the unsupervised hierarchical clustering analysis were selected using significance analysis of microarray (SAM)52 and chi-square analysis, with a 1000-permutation adjustment. The method of the nearest shrunken centroid identified a subgroup of 30 overexpressed (Table S2) and 50 underexpressed (Table S3) genes unique to each subgroup, and these genes were then used to develop a class predictor using prediction analysis for microarrays (PAMs) in R version 2.1.1 (PAM, Stanford, CA).53 The prediction error was calculated by means of 10-fold cross-validation within the training set followed by use in the test set. Supervised cluster analysis of known classes was performed using GeneCluster2 (Broad Institute, Cambridge, MA).54

An expression-based proliferation index (PI) was calculated using the normalized value of 11 genes associated with proliferation (TOP2A, BIRC3, CCNB2, NEK2, ANAPC7, STK6, BUB1, CDC2, C10orf53, ASPM, and CDC42) scaled to the maximum value among 22 plasma cell samples from 22 healthy donors (defined PI = 1), 414 newly diagnosed myelomas, and 45 myeloma cell lines.55,56 The normalized subgroup PI was shown the relative alteration to NPC by adjusting NPC PI as 1. The one-way ANOVA test for difference of PI across the groups was used.

Survival distributions were presented with the use of the Kaplan-Meier method and compared with the log-rank test. Statistical tests were performed with the software package SPSS 12.0 (SPSS, Chicago, IL).

**Results**

**Identification and validation of 7 subgroups in newly diagnosed myeloma based on common gene expression signatures**

Unsupervised hierarchic cluster analysis produced 2 major dendrogram branches with 7 subbranches, which in turn were strongly influenced by the coordinated overexpression of specific genes, many with anchoring genes such as c-MAF and MAFB, CCND3, ASS, IL6R, MMSET, FGFR3, CCNB2, FRZB, and DKK1 (Figure 1).

Application of the PAM model to the training set classified 98% of the samples correctly based on the original unsupervised hierarchic clustering subgroup designations (Table 1). A colorogram of the expression levels of the 700 PAM genes across the training cohort provides a visual reference of the unique gene expression patterns distinguishing the 7 subgroups (Figure 2A). Application of the PAM model to an independent test set produced a resultant colorogram of the 7 classes and the expression levels of the 700-classifier genes (Figure 2B). Although training and test sets contained a different total number of cases, their proportions in each of the 7 subgroups were comparable in training and test sets (Table S4).

**Genetic signatures of expression-defined subgroups**

Translocations between the immunoglobulin heavy chain locus and CCND1, CCND3, c-MAF, MAFB, FGFR3, and MMSET represent...
The reciprocal t(4;14)(p16;q32) translocation results in the hyperactivation of both the FGFR3 and MMSET genes. The majority of cases with spiked FGFR3 or MMSET expression clustered together in one subgroup in both training (Figure 3A) and test (Figure 3B) sets. Of importance, 25% of these cases exhibited only a MMSET spike. Conversely, loss of MMSET expression in FGFR3-positive tumors was not observed. Thus, consistent with a central role of MMSET in driving downstream transcriptional events, cases with MMSET spikes but lacking FGFR3 spikes clustered together with samples exhibiting activation of both genes (Figure 3A-B). Since the MMSET spike represents a dominant feature of group 3, this group was designated as the MS (MMSET) group. While FGFR3 and WHSC1/MMSET represented the top-ranked overexpressed genes in the MS group, other notable genes included the cadherin family member, desmoglein 2 (DSG2), Wnt receptors FZD2 and FZD8, and the B-cell oncogene PBX1. Significant underexpressed genes of potential relevance included the adhesion molecules ICAM4, N-cadherin (CDH2), cadherin 7 (CDH7), and the B-cell differentiation transcription factor PAX5.

Two cyclin D family members are activated by translocations in MM: cyclin D1 by the t(11;14)(q13;q32) in 17% and CCND3 by t(6;14)(p21;q32) in 2%. As with MAF and MAFB spikes, cases with CCND1 and CCND3 spikes clustered together in the training (Figure 3A) and test (Figure 3B) sets, suggesting that activation of 2 cyclin D orthologs results in dysregulation of common downstream transcriptional programs (Figure 3). Unlike MAF and MAFB and FGFR3 and MMSET spikes, which together comprised

Table 1. Confusion matrix of subgroup designations by unsupervised hierarchic clustering and the PAM model in the training set

<table>
<thead>
<tr>
<th>PAM designation</th>
<th>G1</th>
<th>G2</th>
<th>G3</th>
<th>G4</th>
<th>G5</th>
<th>G6</th>
<th>G7</th>
<th>Class error rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>PR</td>
<td>29</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0.048</td>
</tr>
<tr>
<td>LB</td>
<td>1</td>
<td>28</td>
<td>0</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0.011</td>
</tr>
<tr>
<td>MS</td>
<td>0</td>
<td>0</td>
<td>44</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0.048</td>
</tr>
<tr>
<td>HY</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>66</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0.027</td>
</tr>
<tr>
<td>CD-1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>20</td>
<td>1</td>
<td>0</td>
<td>0.091</td>
</tr>
<tr>
<td>CD-2</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>43</td>
<td>0</td>
<td>0</td>
<td>0.001</td>
</tr>
<tr>
<td>MF</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>20</td>
<td>0.048</td>
</tr>
</tbody>
</table>

The t(14;16)(q32;q23) and t(14;20)(q32;q11) translocations result in activation of c-MAF and MAFB proto-oncogenes, respectively, and are together seen in approximately 6% of cases. Although mutually exclusive, MAF and MAFB spikes clustered together in group 7 (Figure 3A-B), suggesting that ectopic expression of the MAF family of transcription factors results in dysregulation of common downstream targets, justifying an MF (MAF/MAFB) subgroup designation. It is noteworthy that 3 cases within the MF subgroup lacked c-MAF or MAFB spikes, suggesting that other MAF family genes may be activated in these cases. Hurt et al50 have reported that CCND2, CX3CR1, and ITGB7 are targets of the c-MAF transcription factor. Indeed, SAM analysis revealed that CX3CR1 and ITGB7 were among the top 50 overexpressed genes unique to the MF group. CCND2, not on the SAM list, was also expressed in other subgroups, although its expression was highest in the MF subgroup (Figure 3A-B). Additional genes with high SAM scores uniquely overexpressed in the MF group and representing known and putative targets of these transcription factors included the recently identified large MAF family target NUAK1/ARK5 as well as NTRK2, ARID5A, SMARCA1, TLR4, SPP1, and G6MB6. The Wnt signaling antagonist SFRP2 was also uniquely overexpressed in this group. Of the SAM-defined underexpressed genes, TNF-induced gene, TNFAP8, was the most significant. Another notable underexpressed gene in the MF group was DKK1, the overexpression of which has been implicated in MM-related bone disease; indeed, the MF group as a whole exhibited a relatively low incidence of bone lesions (Figure 3).
Figure 3. Subgroups are characterized by unique expression patterns. The Affymetrix signal (expression level: vertical axis) of MAF, MAFB, FGFR3, MMSET, CCND1, CCND2, CCND3, FRZB, and DKK1 from the 256 and 158 cases based on the clustergram sample distribution from Figure 2A and B, respectively. The expression levels for each gene are proportional to the height of each bar (representing a single patient sample). Note that spiked expression of CCND1, MAF, and MAFB, and FGFR3 and MMSET is strongly correlated with specific subgroup designations. Also note that cases retaining the MMSET spike but lacking FGFR3 spikes maintain similar cluster designation, and MAF and MAFB spikes cluster in the same subgroups. Several MMSET spike–positive cases in the proliferation subgroup. CCND2 expression was mutually exclusive of CCND1 expression. While highly correlated with the hyperdiploid subgroup, FRZB and DKK1 were both significantly underexpressed in groups LB and MF.

Table 2. Percentages of hyperdiploidy and nonhyperdiploid karyotypes in training (TR) and test (TE) sets

<table>
<thead>
<tr>
<th></th>
<th>PR</th>
<th>LB</th>
<th>MS</th>
<th>HY</th>
<th>CD-1</th>
<th>CD-2</th>
<th>MF</th>
</tr>
</thead>
<tbody>
<tr>
<td>TT2 Hyperdiploid</td>
<td>55</td>
<td>67</td>
<td>43</td>
<td>93</td>
<td>20</td>
<td>20</td>
<td>25</td>
</tr>
<tr>
<td>Nonhyperdiploid</td>
<td>45</td>
<td>33</td>
<td>57</td>
<td>7</td>
<td>80</td>
<td>80</td>
<td>75</td>
</tr>
<tr>
<td>TT3 Hyperdiploid</td>
<td>53</td>
<td>25</td>
<td>33</td>
<td>86</td>
<td>0</td>
<td>50</td>
<td>17</td>
</tr>
<tr>
<td>Nonhyperdiploid</td>
<td>47</td>
<td>75</td>
<td>67</td>
<td>14</td>
<td>100</td>
<td>50</td>
<td>83</td>
</tr>
</tbody>
</table>

Values indicate the percentage of the total number of cytogenetics abnormality cases within each subgroup having the variable indicated. 

$P < .001$ for both TT2 and TT3 groups.
gene BIK, HES5, HIF1A, and SMAD1. In contrast to the HY group, interferon-induced genes IFI127, IFI35, IFIT5, STAT1, and STAT2 were underexpressed in this group. As in MS and MF groups, group 2 expressed relatively high levels of the IL6LR and low levels of the WNT signaling antagonists RFRB (P < .001) and DKK1 (P < .001) relative to the other groups. Overexpression of these latter genes has been linked to the presence of focal bone disease and osteolytic lesions. Consistent with the low expression of DKK1, group 2 had a significantly lower number of magnetic resonance imaging (MRI)–defined focal lesions than seen in the other groups in both the training and test cohorts (Table 2). In lieu of any clear genetic signatures distinguishing this group, we termed it the low bone disease (LB) group.

Group 1 of the unsupervised clustering dendrogram was characterized by the overexpression of numerous cell cycle– and proliferation-related genes (eg, CCNB1, CCNB2, MCM2, CDC2, BUB1, CDC2, TYMS) and cancer–testis antigen genes (eg, MAGE6, MAGEA3, GAGE1, GAGE4). This group also had a significantly higher gene expression–defined proliferation index (PI) than the other groups in both training and test sets, justifying its designation as proliferation (PR) subgroup. All the MM subgroups defined here had a higher PI than plasma cells from healthy donors. In addition, the PR group had a PI similar to that of human MM cell lines (P < .001) (data not shown). Metaphase cytogenetic abnormalities were present in an extraordinarily high 69% of cases in the training group and 83% in the test group (the mean for the remaining newly diagnosed cases was approximately 20%). Both hyperdiploid and nonhyperdiploid cases were equally common, with and without concomitant spikes (Table 2). The training and test sets contained 6 MMSET, 3 CCND1, and 2 MAF spikes. Consistent with the emergence of a PR signature overtime, a number of diagnostic samples in virtually all subgroups, especially in the MF group, exhibited overexpressed subsets of genes defining the PR subgroup (Figure 3A–B).

**Cyclin D expression in subgroups**

Dysregulated expression of 1 of the 3 cyclin D genes (CCND1, CCND2, or CCND3) is a feature of virtually every case of newly diagnosed MM. Hyperactivated expression of CCND1 and CCND2 was seen in more than 95% of the cases studied here, but relative levels and distribution varied across the subgroups in both training and test sets. Expression of 1 of the 3 cyclin D genes is mutually exclusive, and a trend was noted for CCND2 to be expressed in the PR, LB, and MS groups (an occasional CCND1 spike in the PR group may reflect MM progression) and was expressed at highest levels in the MF group. As expected, ectopic low-level expression of CCND1 was observed in the HY group. CCND1 and CCND3 genes exhibited mutually exclusive spiked expression in the CD-1 and CD-2 groups (except for one CCND3 case in the training set clustering in the HY group).

**Laboratory characteristics in the 7 subgroups in the training and test cohorts**

Significant differences in standard laboratory features were noted across subgroups (Table 3): the PR group had a higher incidence of abnormal cytogenetics (P < .001), higher serum levels of B2M (P < .001) and LDH (P < .001), and lower levels of albumin (P < .033). Low albumin and high B2M levels were also observed in MS and MF groups, respectively. A striking difference across subgroups was the lower incidence of MRI-defined bone lesions in LB (30% in the training set and 21% in test set) compared with the remaining groups (P < .01). Thus, gene expression patterns and clinical parameters are highly correlated; the higher incidence of high-risk variables in the PR and MS groups is consistent with their poor prognosis (Figure 4, Table 3).

**Event-free survival and overall survival differ in the 7 subgroups, and molecular class designation is an independent predictor on multivariate analysis**

With a 36-month median follow-up on the training cohort, HY, CD-1, CD-2, and LB subgroups were associated with superior event-free (Figure 4A) and overall (Figure 4B) survival relative to the PR, MS, and MF groups. Kaplan-Meier plots suggested a natural cut between high-risk and low-risk diseases, with CD-1, CD-2, LB, and HY groups representing low risk and MF, MS, and PR cases, high risk (Figure 4B), with 48-month estimates of event-free survival of 68% versus 31% (P < .001) and of overall survival of 79% versus 51% (P < .001), respectively. On multivariate analysis, these genetic groups, along with abnormal cytogenetics and elevation of serum levels of B2M and LDH, were significant independent predictors of survival (Table 4).

**Chromosome distribution of SAM-defined overexpressed genes**

The chromosome map positions of all SAM-defined overexpressed genes for each of the 7 subgroups were determined. In the case of hyperdiploid MM, a significantly higher number of overexpressed genes mapped to chromosomes 3, 5, 7, 9, 11, 15, and 19 (data not shown). The number of overexpressed genes mapping to chromosome 1q was significantly higher in the poor-risk PR and MS groups (Table 5); the number of overexpressed genes mapping to chromosome 1p was significantly higher in the CD-2 group (Table 5).

| Table 3. Correlation of clinical features across GEP-defined subgroups in training (TR) and test (TE) cohorts |
|---|---|---|---|---|---|---|---|---|---|---|---|---|---|
| PR | TR | TE | P | TR | TE | P | TR | TE | P | TR | TE | P |
| B2M level of 339 nM or higher | 66 | 61 | NS | 35 | 41 | NS | 30 | 29 | 17 | 42 | 41 | NS | 52 | 44 | < .01 |
| LDH level of 190 U/L or higher | 48 | 56 | NS | 29 | 22 | NS | 26 | 38 | 17 | 40 | 29 | NS | 45 | 38 | < .01 |
| Albumin level of less than 35 g/L | 66 | 61 | NS | 35 | 41 | NS | 30 | 29 | 17 | 42 | 41 | NS | 52 | 44 | < .01 |
| LDH level of 190 U/L or higher | 48 | 56 | NS | 29 | 22 | NS | 26 | 38 | 17 | 40 | 29 | NS | 45 | 38 | < .01 |
| Albumin level of less than 35 g/L | 66 | 61 | NS | 35 | 41 | NS | 30 | 29 | 17 | 42 | 41 | NS | 52 | 44 | < .01 |

For the MS, HY, and MF groups, the P values were not significant; for the CD-1 group, the P values were not applicable. The total P represents the test for differences in the variable across the subgroups. This was performed on the mean of training and test set percentages. The total P was less than .01 for all groups. Values indicate the percentage of the total number of cases within each subgroup having the variable indicated. Statistical tests were performed for differences in the percentage of cases between the training and test sets.

NS indicates not significant.

*Due to small number of CD-1 cases (n = 6) in the TT3 cohort, a statistical analysis was not performed. Differences were established by chi-square test.
Myeloid gene signature in CD138-selected cells is associated with a good prognosis

As recognized here, and in previous microarray analyses of CD138-selected plasma cells, a myeloid gene expression signature was detectable in a substantial proportion of cases in both training and test sets. This signature was sufficiently strong to confound unsupervised hierarchic cluster analyses, and these cases were removed prior to unsupervised hierarchic cluster analysis. A comparison revealed that the 95 cases in the training set with a myeloid signature displayed more favorable baseline features and better survival than the 256 cases lacking this signature. In this excluded group, abnormal cytogenetics, IgA subtype, and higher levels of bone marrow plasmacytosis as well as serum creatinine and B2M were all significantly less common (P < .05). Moreover, event-free survival (P = .017) and overall survival (P = .046) were superior in the excluded groups. Higher levels of bone marrow plasmacytosis and B2M were also significantly less frequent in the myeloid signature-positive subgroup of the test set (P < .05). However, there was no difference between the retained and excluded cases with respect to albumin, LDH, and number of MRI-defined focal lesions in either the training or test sets.

Discussion

Although presenting with the same histologic diagnosis, MM displays an enormous genomic complexity as well as marked variation in clinical characteristics and patient survival. For treatment advances to occur, clinical outcome data have to be interpreted within the framework of genetic entities, as has been proven useful in leukemia and lymphoma. Here, we have provided a comprehensive and integrated view of the myeloma transcriptome in highly enriched tumor cells from a large cohort of newly diagnosed patients. Based on concordant gene expression signatures, predominantly driven by recurrent translocations and hyperdiploidy, MM could be characterized as belonging to 7 distinct

Table 4. Multivariate proportional hazards analysis

<table>
<thead>
<tr>
<th></th>
<th>Event-free survival</th>
<th>Survival</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HR</td>
<td>P</td>
</tr>
<tr>
<td>PR, MS, and MF versus other</td>
<td>37</td>
<td>2.41</td>
</tr>
<tr>
<td>Any cytogenetic abnormalities</td>
<td>38</td>
<td>1.60</td>
</tr>
<tr>
<td>β2-microglobulin level of 339 nm or higher</td>
<td>44</td>
<td>1.47</td>
</tr>
<tr>
<td>LDH level of 190 μL or higher</td>
<td>34</td>
<td>1.69</td>
</tr>
</tbody>
</table>

n = 255 (of 256 patients); 1 patient is missing cytogenetics. There were 115 progression events and 74 deaths. The following variables were not significant: (1) with the sample of 255, age of 65 years or older was not significant in either analysis (P = .07 for EFS); (2) C-reactive protein level of 4 or higher; and (3) albumin level of 3.5 g/dL or less.

*This column refers to the percentage of 255 cases having the incidence of each clinic feature specified in the first column of each row.

Table 5. Number of SAM-defined overexpressed genes mapping to chromosome 1q and 1p in the 7 subgroups

<table>
<thead>
<tr>
<th>Chromosome</th>
<th>Total</th>
<th>PR</th>
<th>LB</th>
<th>MS</th>
<th>HY</th>
<th>CD-1</th>
<th>CD-2</th>
<th>MF</th>
<th>P</th>
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</thead>
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<tr>
<td>1q</td>
<td>2003</td>
<td>178</td>
<td>52</td>
<td>162</td>
<td>22</td>
<td>21</td>
<td>37</td>
<td>38</td>
<td>&lt;.001</td>
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<tr>
<td>1p</td>
<td>2453</td>
<td>39</td>
<td>20</td>
<td>69</td>
<td>67</td>
<td>38</td>
<td>132</td>
<td>46</td>
<td>&lt;.001</td>
</tr>
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</table>
molecular entities. HY, CD-1, CD-2, and LB subgroups were associated with superior event-free and overall survival following high-dose therapy and stem cell transplantation.

Although the molecular classification of newly diagnosed disease presented here was validated, the associations between classes and survival are likely to be dependent on the type of therapy used. It is our belief that the relationship between subgroups and survival should form the basis for modification and continual evolution of therapies toward subgroup-specific trials. While many of the subgroups defined in this paper are doing extraordinarily well on TT2, the MS and PR groups do not appear to benefit from this therapeutic strategy. However, future therapies that might exploit molecular insights presented here should lead to an improvement in outcome for patients with these types of disease. Indeed, while there is no significant difference in the EFS and OS of 6 of the 7 groups treated on TT3 relative to TT2, the MS group has a significantly longer EFS (P = .04) and trend for better OS (P = .06) on TT3 relative to TT2 (J.D.S., unpublished data, 2006). This benefit will have to be confirmed with longer follow-up of the test set.

When viewed in the context of our previous gene expression classification studies (4 subgroups among 74 cases applying a first generation microarray with only 8000 gene features,22 the validated classification system presented here, with 7 rather than 4 groups, can be explained by an increase both in gene and sample number, enabling the distinction of rare entities (eg, MF, CD-1, CD-2, and PR groups) with very distinct GEP signatures. In a supervised classification system presented here, with 7 rather than 4 groups, the presence of a myeloid gene signature appears to hold important novel information without imposing any predetermined structure on the data; thus, PR and LB groups and the differentiation of the CD-1 and CD-2 groups were identified.

A significant proportion (27%) of newly diagnosed myeloma could not be analyzed for a myeloma signature due to an overwhelming myeloid/normal plasma cell gene expression signature in the post–CD138-selected cells. Postselection flow cytometry clearly showed that while the cases lacking a myeloid gene expression signature were predominantly CD38+/CD45−, the cases with a myeloid gene signature contained both CD38+/CD45+ and CD38+/CD45dim cells (data not shown). The presence of this myeloid cell gene expression signature in CD138-selected cells from healthy donors13 suggests that this reflects a copurification of myeloid cells rather than an aberrant expression of myeloid genes in malignant plasma cells. As myeloid cells do not express CD138, one possible explanation for this finding is that the anti-CD138 antibody binds to immunoglobulin Fc receptors that are highly expressed on cells of the myeloid lineage, which are then collected in the separation procedure. The patients with a myeloid expression signature in the CD138-selected fractions often presented with macrofocal bone marrow involvement with low or even absent diffuse infiltration, as in subjects with MGUS (data not shown). The plasma cell yield in randomly obtained samples depends on whether focal lesions were targeted. CT-guided fine-needle biopsies of focal lesions, recognized on magnetic resonance imaging examinations, will help distinguish whether enriched plasma cells express signatures of the 7 subgroups described here or whether this type of myeloma constitutes a novel subclass. Nevertheless, the presence of a myeloid gene signature appears to hold important clinical information. In addition to lower levels of bone marrow plasmacytosis, this group was associated less frequently with cytogenetic abnormalities and elevations of B2M and creatinine, and most importantly enjoyed superior EFS and OS than the groups lacking this feature. In view of its macrofocal disease presentation, it is noteworthy that the total number of MRI-defined focal lesions did not exceed those cases lacking the myeloid signature. While not appropriate for molecular profiling of the MM transcriptome, the presence of a myeloid gene signature in CD138-selected cells from newly diagnosed MM would denote a favorable disease course.

Several interesting features of the molecular subgroups defined here are worth noting. Relative to the other 5 groups, genes uniquely underexpressed in both CD-1 and CD-2 subgroups included IL6R, HOXB7, BMPR1A, the mitotic cyclin, CCNE, and the cyclin-dependent kinase, CDK6, which has recently been shown to uniquely interact with cyclin D2 in MM plasma cells.61 In a comparison of differentially expressed genes in CD-1 and CD-2 groups, CD-2 was characterized by overexpression of TNFRSF7 (CD27), the SDF-1 receptor CXCR4, CD20, BTG2, and CD38, whereas ASS, INHBE, the proto-oncogene FYN, NID2, and SET7, a gene with homology to MMSET, were overexpressed in CD-1. The biologic and clinical relevance of CD-1 and CD-2 groups is currently unclear, as there was no difference in clinical parameters or survival between the 2 groups. However, CD-2, but not CD-1 was associated with elevated expression of CD20, which has previously been shown to be associated with the t(11;14)(q13; q32)61 and highly correlated with CD20 protein expression in MM plasma cells.42,62 These tumors also expressed other markers of more immature B cells including PAX5 and the surrogate immunoglobulin light chain VPREB. Another striking genetic difference between CD-1 and CD-2 was the significant difference in the elevated expression of genes from the p arm, but not the q arm, of chromosome 1 in CD-2.

Hyperdiploidy is a distinct genetic entity with a good prognosis and largely devoid of common recurrent immunoglobulin-mediated translocations.17-19 The HY MM subgroup was mainly characterized by overexpressed genes derived from the odd number chromosomes 3, 5, 7, 9, 11, 15, 19, and 21; however, this signature was also observed in cases not showing hyperdiploidy by flow cytometry (data not shown); such diploid and hypodiploid cases may be derived through a similar genetic mechanism (trisomies of odd chromosomes), although clonal evolution may result in loss of DNA on other chromosomes so that the DNA complement is essentially diploid. The assignment of both hyperdiploid and nonhyperdiploid cases to the PR group suggests that a simple recognition of hyperdiploidy is insufficient for proper risk assessment: those with a proliferation signature and hyperdiploidy would be at higher risk than those with an HY signature alone.

An important question concerns the influence of various types of genetic insults in the etiology of MM and their subsequent effects on the transcriptome. Using high-resolution array comparative genomic hybridization (aCGH), mRNA microarray, interphase fluorescence in situ hybridization (FISH), and novel bioinformatics approaches, we recently identified 4 different MM subtypes based on recurrent DNA copy number changes.63 Using gene expression profiling as a surrogate to validate the aCGH-defined groups, we were able to confirm the existence of 2 forms of hyperdiploid disease, one containing gains of 1q, deletion of chromosome 13, and absence of trisomy of chromosome 11.63 In this study, we could distinguish only hyperdiploid MM as a single group without the ability to separate out the 2 unique subtypes identified by aCGH. These subgroups and additional groups are likely to emerge as more sophisticated data mining tools are applied to this large dataset.
A striking feature common to the 2 high-risk groups, MS and PR, was the significant number of overexpressed genes mapping to 1q. While elevated expression of 1q genes is apparent de novo in the MS group, the elevated expression of 1q genes is not apparent in the 5 remaining groups and thus appears to be coincident with the acquisition of a proliferation signature. Major questions emerging from these observations are (1) whether there is a common mechanism by which the 1q genes are activated de novo in the MS group and acquired during disease progression in the other groups, and (2) if there are genes mapping to 1q that contribute to the proliferation signature. Indeed, a central role for chromosome 1q abnormalities in myelomagenesis has been suggested. Tandem duplications and jumping translocations of 1q21 occur frequently in this malignancy,64-66 and gain of 1q is one of the most common abnormalities in MM.67-70 As mentioned above, a form of hyperdiploidy characterized by gains of 1q was found to have a poorer clinical outcome than hyperdiploid disease lacking this feature.63 High-resolution aCGH studies also identified a nonhyperdiploid entity characterized by an ampiclon at 1q21.63 Using correlations of gene expression extremes with survival in the training cohort, we recently found that high-risk disease was linked to overexpression of 1q genes and reduced expression of 1p genes (J.D.S., manuscript in preparation). Recent studies using aCGH71 and interphase FISH72 have revealed that gains/amplification of 1q21 accompany the progression of smoldering to overt MM. We have recently shown that gains/amplifications of 1q21 were linked to inferior survival in patients treated on TT2, and the incidence and magnitude of 1q21 gains increased from diagnosis to relapse in patients on this trial.72 Thus, it appears that gain/amplification of 1q21 may be a key genetic event in MM pathogenesis and progression. Whether abnormalities of 1q are a marker of or contribute to disease progression is not currently clear.

An important unanswered question is whether a majority of, if not all, MM cases will eventually acquire a PR class designation. Support for this concept comes from data derived from applying the PAM model to relapsed cases. While present in approximately 18% of newly diagnosed MM, a PR signature was found in 45% of 83 relapses (J.D.S., unpublished data, November 2005). Further evidence for acquisition of a PR signature during disease progression comes from data showing that while 30 of 35 cases with paired baseline and relapse samples maintain the same class designation at relapse, 1 MS and 4 HY cases at diagnosis shifted to a PR classification at relapse (J.D.S., unpublished data). Overexpression of proliferation-associated genes in cases within each of the 6 other subgroups (Figure 2A-B), presence of spikes in the PR group, and class shift to PR with disease progression all suggest that the acquisition of the PR designation is a feature that will emerge in most if not all relapsing MM cases.

The lack of progress in prolonging survival in patients with high-risk MM, now best identified by molecular tests, and the possibility that all MM will evolve to acquire an aggressive PR-like disease should encourage development of therapeutics that target the molecular pathways unique to high-risk disease elucidated through genomic profiling.

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