Integrative Genomics Identifies Molecular Alterations that Challenge the Linear Model of Melanoma Progression

Amy E. Rose¹, Laura Poliseno¹, Jinhua Wang², Michael Clark¹, Alexander Pearlman², Guimin Wang¹, Eleazar C. Vega y Saenz de Miera¹, Ratna Medicherla¹, Paul J. Christos⁷, Richard Shapiro³,⁶, Anna Pavlick⁴,⁶, Farbod Darvishian⁵,⁶, Jiri Zavadil⁵, David Polsky¹,⁵,⁶, Eva Hernando⁵,⁶, Harry Ostrer²,⁴,⁵,⁶, and Iman Osman¹,⁴,⁶

Abstract

Superficial spreading melanoma (SSM) and nodular melanoma (NM) are believed to represent sequential phases of linear progression from radial to vertical growth. Several lines of clinical, pathologic, and epidemiologic evidence suggest, however, that SSM and NM might be the result of independent pathways of tumor development. We utilized an integrative genomic approach that combines single nucleotide polymorphism array (6.0; Affymetrix) with gene expression array (U133A 2.0; Affymetrix) to examine molecular differences between SSM and NM. Pathway analysis of the most differentially expressed genes between SSM and NM (N = 114) revealed significant differences related to metabolic processes. We identified 8 genes (DIS3, FGFR1OP, G3BP2, GALNT7, MTAP, SEC23IP, USO1, and ZNF668) in which NM/SSM-specific copy number alterations correlated with differential gene expression (P < 0.05; Spearman's rank). SSM-specific genomic deletions in G3BP2, MTAP, and SEC23IP were independently verified in two external data sets. Forced overexpression of metabolism-related gene MTAP (methylthioadenosine phosphorylase) in SSM resulted in reduced cell growth. The differential expression of another metabolic-related gene, aldehyde dehydrogenase 7A1 (ALDH7A1), was validated at the protein level by using tissue microarrays of human melanoma. In addition, we show that the decreased ALDH7A1 expression in SSM may be the result of epigenetic modifications. Our data reveal recurrent genomic deletions in SSM not present in NM, which challenge the linear model of melanoma progression. Furthermore, our data suggest a role for altered regulation of metabolism-related genes as a possible cause of the different clinical behavior of SSM and NM. Cancer Res; 71(7); 2561–71. ©2011 AACR

Introduction

Superficial spreading melanoma (SSM) and nodular melanoma (NM), the 2 most common histopathologic subtypes (70% and 20% respectively), are characterized by markedly different clinical presentations and natural histories. NM has a higher rate of recurrence (1, 2) and has not shown the same degree of downward-stage migration at initial presentation relative to SSM (3). These differences are generally attributed solely to the advanced thickness of NM, with no prognostic relevance assigned to histopathologic subtype in melanomas of equivalent thickness. Previous studies by several groups including ours, however, suggest that underlying molecular differences between these 2 subtypes may also contribute to the disparate outcomes (4–6).

It is generally accepted that SSM and NM develop along a linear pathway of progression that begins with transformation of epidermal melanocytes and differs between subtypes primarily with respect to the speed with which the transformed melanocytes invade the dermis (7, 8). Until the identification of distinct molecular alterations characterizing acral melanoma such as c-Kit mutation and amplification (9–11), all melanoma subtypes were typically viewed as a relatively homogenous biological entity (8). Advances in genomic technology in the setting of a broader recognition of the biological heterogeneity of cancer have changed this long-held view, at least, for acral melanoma. As such, clinical trials evaluating the efficacy of c-Kit inhibitor imatinib for patients with metastatic melanoma from an acral primary are ongoing (12). Currently, there is no definitive evidence to show that SSM and NM are molecularly different enough to warrant a unique molecular classification such as the one that has now been assigned to acral melanoma.

It is difficult to reconcile observed differences between SSM and NM, such as the minimal degree of epidermal involvement and the lack of a detectable radial growth phase (RGP) in NM,
with the current linear progression model. To our knowledge, there are no studies that have evaluated the genomic and gene expression alterations that characterize SSM and NM. We examined by an integrative genomic approach the hypothesis that SSM and NM follow separate pathways of development from the transformed melanocyte to the invasive primary melanoma that are characterized by subtype specific molecular alterations.

Materials and Methods

Melanoma cell lines and human tissues

Cell lines included normal melanocytes cultured from infant foreskin (NHM), immortalized melanocytes (Hermes 1 and 2B), primary SSM (WM35, WM1552c, WM1575), primary NM (WM39, WM8532), primary vertical growth phase (VGP; WM98.1, WM8532), metastatic NM (Lu451, SK-147), and metastatic melanoma of unknown primary histologic subtype (501MEL, A375, SK-MEL-19, -29, -85, -94, -100, -103, -173, -187, -192, -197). Metastatic cell lines were kindly provided by Alan Houghton (Memorial Sloan-Kettering Cancer Center, New York), and primary cell lines (and Lu451) were purchased commercially from the Wistar Institute. Cell lines were not further tested or authenticated (see Supplementary Methods for details on cell lines and tissues).

SNP DNA array

DNA was hybridized to the Affymetrix genome-wide human single nucleotide polymorphism (SNP) 6.0 array as per the manufacturer protocol (Affymetrix). CEL files were generated using the GeneChip Command Console software (Affymetrix), and the Birdseed v2 algorithm (13) was used to make the genotype calls and summarize the probe set intensity values for ensuing copy number analysis. A single reference genome profile assayed in the same batch was created using values from the 2 normal melanocytes cultured from infant foreskin (NHM).

Gene expression array

RNA was hybridized to the Affymetrix U133A 2.0 array as per manufacturer protocol. Array data were normalized using the robust multichip average (RMA) and filtered for expression values lower than 16. The first analysis undertaken was an exploratory principal component analysis (PCA), which allows for the visualization of relationships between samples in a multidimensional data set (14). Data were then log transformed to carry out an ANOVA and a 2-sample unpaired t test, with P values adjusted for multiple comparisons (Bonferroni correction). A single-factor, 2-level ANOVA design was utilized to examine sources of measured variability between the SSM and NM. A software package (Partek) was used to facilitate the analysis. MAICE-compliant data for the Affymetrix SNP6.0 and U133A 2.0 gene expression array have been deposited in the National Center for Biotechnology Information Gene Expression Omnibus [GSE22306 (GSE22301 and GSE22305)].

Integrative genomic analyses

Correlation of copy number data with gene expression was done using the Partek Genomic Suite (Partek). Copy number gains and losses were detected using a hidden Markov model algorithm in the Partek copy number workflow for unpaired samples (see Supplementary Methods for detailed analysis).

External validation

Unprocessed data for 13SSM and 5NM human tissues were downloaded from the 2007 publication by Jaeger and colleagues (4). Data were then normalized using RMA and log transformed for a 2-sample unpaired t test with a Bonferroni correction for multiple testing. GenePattern was used for clustering and gene set enrichment analysis. The complete list of 540 transcripts differentially expressed between RGP and VGP in the 2009 publication by Scatolini and colleagues (15) was also used as an external validation.

Quantitative reverse transcriptase PCR

Quantitative reverse transcriptase PCR (qRT-PCR) was used to verify the results of gene expression microarray data. Data were exported using SDS 2.3 and analyzed using the methods of Pfaffl (16), assuming 100% efficiency of amplification with normalization to 4 housekeeping genes: TMPS52, PDLIM4, DHX15, and SNF8. These genes were selected on the basis of their uniform expression across the data set (coefficient of variance <10%). Fold change was evaluated relative to melanocyte controls. Primer sequences are summarized in Supplementary Table S1A.

Genomic qPCR

Five genes were evaluated for validation, using genomic PCR: D5S3, G3BP2, MTAP, SEC23IP, and USO1. Gene-specific primers are shown in Supplementary Table S1B, and housekeeping genes UBE2E1 and GNS were used as previously described (ref. 17; see Supplementary Methods for detailed protocol).

Western blot

Membranes were probed with anti-MTAP (methylthioadenosine phosphorylase) polyclonal antibody (Santa Cruz Biotechnology); anti-ALDH7A1 polyclonal antibody (Epitomics); and anti-EPB41L3 (Abnova) at dilutions of 1:200, 1:1,000, and 1:500, respectively (see Supplementary Methods for detailed protocol).

Lentiviral infection and growth curve

MTAP coding sequence was amplified from pBS/MTAP (MHS1010-7295727; Open Biosystems) by PCR and then cloned into the PetI and Xdel restriction site of pWPI lentiviral vector (gift of Dr. Hernando), downstream of the EFl promoter (see Supplementary Methods for detailed protocol).

Immunohistochemistry on tissue microarrays

Immunohistochemistry (IHC) was done using a tissue microarray (TMA) of 20NM and 20SSM cases. Tissue cores were evaluated for expression of ALDH7A1 (rabbit polyclonal antibody, Abcam Inc.; catalogue #ab80187) diluted 1:500,
EPB41L3 (purified MaxPab mouse polyclonal antibody; Abnova Corporation, catalogue #H00023136-B01P) at 1:500, and MTAP [monoclonal antibody (M01) clone 2G4 purified mouse immunoglobulin; Abnova; catalogue #H00004507-M01] at 1:2,000 (see Supplementary Methods for detailed protocol).

An attending pathologist (F.D.) scored the expression of ALDH7A1, MTAP, and EPB41L3 in each core on a scale from 0 to 2. The association between staining intensity and histologic subtype was assessed by the t test or χ² test for each marker. The χ² test was also used to compare the number of cores with a specific intensity score between the NM and SSM groups. A multivariate, general, linear model was used to compare the adjusted mean expression of ALDH7A1 between the NM and SSM case groups after controlling for thickness and ulceration. All P values were 2-sided, with statistical significance evaluated at the 0.05 α level. All analyses were done in SPSS version 18.0 (SPSS Inc.).

Analysis of genetic and epigenetic alterations of ALDH7A1

ALDH7A1 exons 2, 12, 14, and 15 and introns 5 and 16 were sequenced (Supplementary Methods, primers shown in Supplementary Table S1C). To analyze the methylation and acetylation status of ALDH7A1 promoter, WM1552c cells were treated with 1 μmol/L of methyltransferase inhibitor 5-aza-cytidine (aza-CR), followed by an additional 24-hour incubation with 1 μmol/L aza-CR plus 200 ng/mL of histone deacetylase inhibitor trichostatin A (TSA). ALDH7A1 mRNA expression level was analyzed by real-time PCR (primers shown in Supplementary Table S1D; see Supplementary Methods for detailed protocol).

Results

SNP array identifies recurrent NM/SSM-specific copy number alterations

To investigate the genomic alterations characteristic of SSM versus NM, we conducted high-resolution SNP array on a panel of 22 melanoma cell lines inclusive of normal melanocytes, primary SSM (WM1552c, WM35), primary NM (WM39, WM278), metastatic NM (Lu451, SK-MEL-147), and metastatic melanoma. The list of significantly altered genomic segments across all the cell lines was then filtered to identify those specific to SSM and NM. SNP array (Supplementary Table S2) identified 408 SSM-specific regions with significant genomic alteration (212 gains, 196 losses) and 543 NM-specific genomic alterations (295 gains, 248 losses).

Gene expression profiling reveals 114 genes differentially expressed between SSM and NM

PCA showed that the melanocytes formed a tight and distinct cluster, whereas there was a high degree of variance among the melanoma cell lines. SSM cell lines clustered together and seemed to be closely related to a larger group of metastatic cell lines. On the contrary, the NM cell lines, particularly the NM primaries, did not cluster together and also did not fall within the larger cluster of metastatic and SSM cell lines. All but 1 of the 4 NMs fell outside 2 SDs of the mean expression value for all other melanomas (including SSM and metastatic). Both of the NM primaries showed distinct expression profiles relative to the other melanoma cell lines (data not shown). These results, although exploratory in nature, suggested that there was enough difference with regard to the gene expression profiles between NM and SSM to justify further investigation of the sources of these differences.

Two statistical analyses were then done to identify differentially expressed genes between SSM and NM. The first analysis (ANOVA) revealed a list of 79 gene probe IDs differentially expressed between SSM and NM relative to each other and relative to normal melanocytes and metastatic melanoma cells (P < 0.05; Bonferroni correction; Fig. 1A). The second statistical analysis (t test; SSM vs. NM) identified 114 differentially expressed genes (116 probe IDs; P < 0.05; Bonferroni correction; Fig. 1B and Supplementary Table S3).

Functional annotation shows that differentially expressed transcripts between SSM and NM are related to metabolic pathways

Gene ontology (GO) analysis of differentially expressed genes between SSM and NM revealed enrichment for GO biological process (BP) terms related to primary, cellular, and nucleic acid metabolic processes (Supplementary Table S4). The most enriched term with regard to the number of genes was "metabolic process" (total gene count 65, 55.1%; P = 0.04), which included genes such as MTAP, ALDH7A1, angiogenin (ANG), and dihydrofolate reductase (DHFR; Supplementary Table S4). The 2 most statistically significant GO BP terms were "nucleic acid metabolic process" (total gene count 38, 32.2%; P = 0.003) and "cellular component organization/biogenesis" (total gene count 27, 22.9%; P = 0.001).

To prioritize the 114 differentially expressed NM/SSM, we focused on the 25 genes with differential expression in both analyses (the ANOVA and t test; Supplementary Table S5). The gene showing the highest and most significant differential expression between SSM and NM was MTAP (fold change = −21.5; P = 0.0003), in which expression was lower in SSM. The gene with the second highest fold change between NM and SSM expression was ALDH7A1 (fold change = −18.0; P = 0.03), and the second most significant difference was erythrocyte membrane protein band 4.1 (EPB41L3, fold change = −6.2; P = 0.001; Supplementary Table S5). MTAP, ALDH7A1, and EPB41L3 were underepressed in SSM relative to NM, and all genes were related to cellular metabolism, consistent with the results of the pathway analysis (Supplementary Table S4).

Integrative genomic workflow yields 8 genes in which SSM/NM-specific copy number changes are correlated with NM/SSM differential expression

NM/SSM-specific copy number gains and losses that mapped to regions coding for genes found to be differentially expressed between NM and SSM in our initial analysis (N = 114) were evaluated for statistical correlation between copy number and mRNA expression level. The integrative analysis revealed a list of 8 genes (BDBC, FGFR1OP, G3BP2, GALNT7, MTAP, SEC23IP, USO1, and ZNF668) that met the following
criteria: (i) NM/SSM-specific copy number gain or loss; (ii) differential gene expression between NM and SSM; and (iii) copy number significantly correlated with gene expression (Table 1). Included in this list of 8 genes were MTAP and FGFR1OP, the 2 genes with the highest degree of fold change between NM and SSM in opposite directions (MTAP lower in SSM, FGFR1OP higher in SSM), suggesting that these genes may hold increased biological relevance.

### Table 1

<table>
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<tr>
<th>Gene symbol</th>
<th>Gene expression</th>
<th>P-value&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Copy number status</th>
<th>Correlation&lt;sup&gt;b&lt;/sup&gt;</th>
<th>P-value&lt;sup&gt;c&lt;/sup&gt;</th>
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<tr>
<td>FGFR1OP</td>
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<td>DEL NM</td>
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<td>0.005</td>
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<td>DEL SSM</td>
<td>0.94</td>
<td>0.005</td>
</tr>
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<td>SSM-NM</td>
<td>0.009</td>
<td>AMP SSM</td>
<td>0.94</td>
<td>0.005</td>
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<td>0.005</td>
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<tr>
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<td>DEL SSM</td>
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<td>0.04</td>
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<tr>
<td>ZNF668</td>
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<td>DEL NM</td>
<td>0.83</td>
<td>0.04</td>
</tr>
<tr>
<td>USO1</td>
<td>SSM-NM</td>
<td>0.04</td>
<td>DEL SSM</td>
<td>0.83</td>
<td>0.04</td>
</tr>
</tbody>
</table>

<sup>a</sup>The t test, with Bonferroni correction for multiple comparisons.

<sup>b</sup>Spearman’s rank correlation coefficient.

<sup>c</sup>By Spearman’s rank test.
Array results verified using genomic qPCR and qRT-PCR

The copy number status of USO1, G3BP2, DIS3, and SEC23IP was evaluated in an expanded panel of NM/VGP cell lines (N = 7) and SSM/RGP cell lines (N = 4). Genomic PCR verified the presence of significant deletions in 100% (P < 0.05; N = 4) of SSM cell lines relative to melanocytes in the regions of USO1, G3BP2, and DIS3 (Fig. 2A, for DIS3). In the region of SEC23IP, genomic PCR verified significant deletions in 75% (P < 0.01; N = 4) of the SSM cell lines (Fig. 2B). Of note, mitotic
control homologue DIS3 was significantly deleted in all 4 RGP-like SSM cell lines assessed by genomic PCR whereas 4 of the 7 VGP-like NM cell lines were diploid and 2 had significant ($P < 0.01$) amplification (Fig. 2A). The same 5 genes were also evaluated for mRNA expression by qRT-PCR, using the melanoma cell lines from the array. qRT-PCR verified the pattern of differential gene expression observed in the array for all genes (i.e., down in SSM relative to NM), with 3 genes (MTAP, DIS3, and SEC23IP) showing a statistically significant ($P < 0.05$; t test) difference in expression between NM and SSM (DIS3 and SEC23IP; Fig. 2; MTAP, Fig. 4C).

Array results verified by *in silico* analysis of external data sets

Reanalysis of a publicly available gene expression data set by Jaeger and colleagues (4) yielded a list of 291 transcripts differentially expressed between NM and SSM (not shown). Also used as an external validation was a data set from the 2009 study by Scatolini and colleagues (15). All 5 genes with NM/SSM-specific copy number and gene expression changes validated using quantitative PCR (qPCR) and qRT-PCR (USO1, G3BP2, DIS3, SEC23IP, and MTAP) were validated as having the same trend of expression (i.e., down in SSM relative to NM) in the external data set by Jaeger and colleagues (DIS3; Fig. 3; not shown). Only 3 of these genes (MTAP, SEC23IP, and G3BP2) were included in the data set of Scatolini and colleagues, but all were validated as having the same trend of expression between subtypes, with G3BP2 also showing statistically significant lower expression in SSM than in NM in the external data set (15).

Overexpression of MTAP in SSM reduces cell growth

The SNP array identified 2 SSM-specific deletions in the region of MTAP (9p21.3) that were significantly correlated with gene expression (Fig. 4A). The deletion in SSM cell line WM35 is more focal, affecting only the last 4 of 8 exons (Fig. 4A), whereas the deletion in SSM cell line WM35 is more focal, affecting only the last 4 of 8 exons (Fig. 4A). Consistent with the array results, genomic PCR of the MTAP gene for cell line WM1552c showed complete loss of DNA in all 3 exons assessed (exons 1, 5, and 8; Fig. 4B). Genomic PCR of MTAP in SSM cell line WM35 showed retained DNA in exon 1 but loss in exons 5 and 8, which is consistent with the position of the copy number loss identified by the SNP array (Fig. 4B). Furthermore, ectopic expression of MTAP by lentiviral infection caused a significant decrease in the growth of SSM cell line WM1552c, supporting an oncosuppressor role (Fig. 4D).

Validation of array results, using Western blot

ALDH7A1 and EPB41L3 were also prioritized for protein validation studies because, second only to MTAP, they had the highest (ALDH7A1) and most significant (EPB41L3) fold changes noted in the gene expression array (Supplementary Table S5). Expression of ALDH7A1, EPB41L3, and MTAP was assessed by Western blotting in the same primary melanoma cell lines used in the array as well as an additional SSM cell line (WM1575) and 2 additional RGP-like cell lines (WM98.1 and WM853.2). Consistent with the array results, ALDH7A1, a gene involved in the detoxification of aldehydes generated by lipid peroxidation, showed complete loss of expression in 2 of the 3 SSM cell lines relative to NM and melanocyte controls (Fig. 5A). Also consistent with the array results, EPB41L3 showed complete loss of expression in all SSM cell lines relative to melanocytes. EPB41L3 showed higher expression in NM cell lines than in SSM but lower expression in NM than in normal, which also verified the array results (Fig. 5A). MTAP expression was completely lost in the 2 SSM cell lines from the array (WM35 and WM1552c), which was consistent with the genomic loss, and showed lower expression in the additional SSM cell line (WM1575) than in both NM and normal melanocytes.
Regulation of ALDH7A1 expression in SSM

We first carried out mutational analysis of ALDH7A1 (exons 2, 12, 14; introns 5, 16) in 7 VGP and 4 RGP cell lines that revealed no evidence of mutation (Supplementary Table S6). We then explored whether the lower expression of ALDH7A1 in SSM is due to the epigenetic mechanisms. We found a significant (P < 0.01) increase in ALDH7A1 mRNA levels in SSM WM1552c cells after treatment with 1 μmol/L aza-CR and 200 ng/mL TSA (Fig. 5B). These data suggest that a combination of hypermethylation and deacetylation may be responsible for the lower expression of ALDH7A1 in SSM.

Validation of array results using IHC

Expression of ALDH7A1, EPB41L3, and MTAP was further evaluated using TMAs of 20 NM and 20 SSM human primary melanoma tissues. Baseline demographic and clinicopathologic data for these cases are presented in Table 2. ALDH7A1 expression was significantly higher in NM cores than in SSM cores. The mean intensity score (on a scale from 0 to 2) of NM cores (N = 20) was 1.62 (SD = 0.43) compared with 1.03 (SD = 0.47) in the SSM cores (N = 20; P < 0.0001; t test). In a multivariate, general, linear model controlling for tumor thickness and ulceration, the adjusted mean staining intensity score remained significantly higher in NM (1.67, SE = 0.11; 95% CI = 1.45–1.88) than in SSM (0.97, SE = 0.11; 95% CI = 0.76–1.19; P < 0.0001). The ALDH7A1-positive cores showed a distinctive, perinuclear “dot-like” intensification consistent with a Golgi pattern of staining (Fig. 5C). Almost all cores that were strongly positive (score 2) for EPB41L3 showed a membranous pattern of staining which was more common in SSM than in NM [46% (N = 18) and 11% (N = 5), respectively; P = 0.001; data not shown]. There was no significant difference noted in the intensity of MTAP staining between NM and SSM.

Discussion

Our results suggest that SSM and NM are not sequential phases of linear progression but rather 2 different biological entities that are characterized by distinct molecular...
alterations. Genomic alterations that showed a significant correlation with gene expression were predominantly SSM-specific deletions. Although there are many posttranscriptional and posttranslational events that can affect levels of gene expression, the identification of recurrent SSM deletions that are retained or even amplified in NM cannot be reconciled with the current paradigm of linear progression.

GO analysis of differentially expressed genes between SSM and NM showed enrichment for genes related to metabolic processes. ALDH7A1 showed higher expression in human NM tissues than in SSM tissues that remained significant in a multivariate model controlling for thickness and ulceration. Recent studies have shown that expression of aldehyde dehydrogenases can identify the presence of cancer stem cell populations in hepatocellular carcinoma and breast cancer (18, 19). Overexpression of ALDH7A1 in NM versus SSM may reflect differences in the percentage of tumor-initiating cells that could explain rapid vertical growth of NM. Not only was expression of ALDH7A1 higher in NM than in SSM and in normal skin but also its expression was lower in SSM than in normal skin. Our data suggest that, in the absence of genomic losses and premature STOP codons, the lower expression of ALDH7A1 in SSM may be due to an epigenetic mechanism.

ALDH7A1 is highly conserved throughout evolution and is homologous to ALDH7B1 in plants, where overexpression of the enzyme is believed to indicate a global upregulation of the organism’s response to environmental stress (20). Similarly, ALDH7 and other ALDH family members are highly expressed in the lens of the murine eye, where they have a protective effect against UV-induced DNA damage (21). Thus, it is possible that the loss of ALDH7A1 observed in SSM makes the subtype more susceptible to and/or more likely to be the result of carcinogenic exposures such as UV light.
The utilization of an integrative genomic approach makes it more likely that the alterations identified are biologically relevant. Genes previously identified using microarrays as differentially expressed between NM and SSM have not been further studied at the functional level; thus, the underlying mechanisms that contribute to differential gene expression are not understood. Of note, one previous gene expression study of RGP and VGP melanoma revealed the unexpected finding that within the same patient, the expression profile of the metastatic melanoma was consistent with a RGP-like signature whereas the nodular primary exhibited a separate, VGP-like expression profile (22). This observation is consistent with our PCA of melanoma cell lines showing that SSM primary cell lines cluster with metastatic lesions whereas all of the primary NMs fall outside this cluster. On the basis of the relatively aggressive clinical behavior of NM relative to SSM, it might be expected that NM would cluster with the metastatic lesions. Although we recognized the exploratory nature of the PCA, the findings from these 2 independent studies showing that gene expression profiles of SSM and metastatic lesions are more closely related than the profiles of nodular primaries support the hypothesis that NM and SSM represent different biological processes.

An array-based integrative genomic workflow and subsequent verification studies identified SSM-specific deletions in genes that were significantly correlated with gene expression. USO1 and G3BP2 are adjacent genes on chromosome 4q21.1. USO1 has been primarily studied in yeast in which it is involved in transport between the endoplasmic reticulum and the Golgi apparatus (23). It is notable that both of our in silico analyses of external data sets confirmed our results showing lower expression of G3BP2 in SSM than in NM (4, 15). G3BP2 is a relatively recently characterized GTPase-activating protein that is overexpressed in breast cancer (24) and that plays a role in the regulation of NFκB signaling (25), a pathway known to be altered in melanoma.

DIS3 has been shown to function in the regulation of proper chromosomal segregation during mitosis, and DIS3 mutant yeast cells have increased sensitivity to microtubule-destabilizing agents such as thiabendazole (26). In this regard, molecular compound library screening studies by our group showed that mebendazole, a microtubule-destabilizing drug, showed subtype-specific efficacy such that SSM cell lines were sensitive but NM cell lines were resistant (5). Further supporting the potential biological relevance of the genes identified by our integrative genomic analysis, 2 independent gene expression profiling studies of colorectal cell lines and human tissues have identified overexpression of DIS3 as high as 38-fold in primary and metastatic tumors relative to normal colonic mucosa (27, 28). Thus, the overexpression and increased DNA copy number identified in the region of DIS3 in our NM cell lines may play a role in its rapid vertical growth and aggressive behavior. The same gene, however, also shows genomic deletion and reduced expression in SSM cell lines. Thus, SSM-specific genomic deletions such as the ones noted in the region of DIS3 may play a mechanistic role in the observed increased sensitivity of SSM cells to microtubule-destabilizing agents.

A previous gene expression profiling study identified SEC23IP as an important mediator of bone formation and metabolism (29). Interestingly, 3 of the top candidate genes from that study (IFT13, SEC23IP, and PPP2R2D) were also found to be among the most differentially expressed between NM and SSM in our study. Although the link between these 2 investigations is not immediately apparent, many pathways known to play a role in carcinogenesis, such as interferon signaling and other cytokines, are also important in bone formation and metabolism (30). SEC23IP has also recently

### Table 2. Baseline demographics and clinico-pathologic variables of the 40 primary melanoma cases included on the SSM and NM tissue microarrays

<table>
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<th>Variable</th>
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<th>NM (N = 20)</th>
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<td>13 (65)</td>
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<td>Female</td>
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<td>Age at diagnosis, y (median)</td>
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<td>Stage, n (%)</td>
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</tr>
<tr>
<td>Extremity</td>
<td>5 (25)</td>
<td>8 (40)</td>
</tr>
<tr>
<td>Axial</td>
<td>13 (65)</td>
<td>9 (45)</td>
</tr>
<tr>
<td>Head and neck</td>
<td>2 (10)</td>
<td>3 (15)</td>
</tr>
<tr>
<td>SLN biopsy positive, n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>4 (20)</td>
<td>5 (25)</td>
</tr>
<tr>
<td>No</td>
<td>16 (80)</td>
<td>15 (75)</td>
</tr>
<tr>
<td>Recurrence, n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>5 (25)</td>
<td>6 (30)</td>
</tr>
<tr>
<td>No</td>
<td>15 (75)</td>
<td>14 (70)</td>
</tr>
<tr>
<td>Status at last follow-up, n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Alive</td>
<td>16 (80)</td>
<td>14 (70)</td>
</tr>
<tr>
<td>Dead</td>
<td>4 (20)</td>
<td>6 (30)</td>
</tr>
</tbody>
</table>

Abbreviation: SLN, sentinel lymph node biopsy.
been identified as a candidate gene for Waardenburg syndrome, a disorder characterized by craniofacial defects and pigment abnormalities, after it was shown that knockdown of sec23p in the Xenopus embryo resulted in impaired migration of neural crest cells (31).

MTAP maps to 9p21.3 and is commonly codetiled with tumor suppressor p16 (CDKN2A; 32). It plays a critical role in the salvage pathway of adenine and methionine and has previously been reported to be lost in several tumors including melanoma (33). Our results show lower expression of MTAP in SSM than in NM, which was significantly correlated with SSM-specific genomic deletions in the region of 9p21.3. MTAP loss in SSM may be particularly relevant, given the recent discovery of MTAP variants that are associated with increased number of nevi and with increased risk of melanoma (34, 35). The role of somatic alterations in MTAP with regard to prognosis and tumorigenesis is less clear, but our results suggest that MTAP deletions may be more common in SSM than in NM. In addition, lentiviral mediated over-expression of MTAP in an SSM cell line with a homozygous deletion resulted in decreased tumor growth compared with MTAP null cells, suggesting that MTAP may function as a tumor suppressor in a subtype-specific fashion. Previous studies have shown that MTAP status may be valuable as a predictor of response to interferon, such that only treatment of patients with MTAP-positive tumors confers a survival advantage compared with those who are untreated (36,37). Thus, whether MTAP deletions differentially affect SSM relative to NM, the MTAP status of a tumor may play a role in the development of a subtype-specific treatment strategy. Because MTAP-deficient cells lose the salvage pathway for purine synthesis, they are sensitive to agents that target the de novo purine synthesis pathway. A recent study that evaluated the efficacy of l-alanosine, an inhibitor of de novo adenine synthesis, for the treatment of MTAP-deficient solid malignancies showed no objective responses (38). The authors suggested that improved patient selection based on more standardized, DNA-based techniques for identifying MTAP loss such as FISH or genomic PCR might improve results of future studies.

In conclusion, our data support a separate molecular classification of SSM and NM. Further functional studies of genes that are specifically deleted in SSM may better define their roles as potential drivers of subtype-specific tumorigenesis.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

Acknowledgments

The authors thank Yutong Zhang of the New York University Cancer Institute and Connie Zhao of the Genomics Resource Center of Rockefeller University for Technical Assistance in carrying out the gene and SNP arrays.

Grant Support

This work was supported by NYU Cancer Institute Cancer Center Support Grant (5 P30 CA016087), the Chemotherapy Foundation, the Skin Cancer Foundation, the Elsa Pardoe Foundation, and the Marc Jacobs Campaign to support melanoma research (I. Osman). The work was partially supported by the Cornell Clinical Translational Science Center (UL1-RR024996) to P.J. Christos.

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Received August 10, 2010; revised December 19, 2010; accepted February 3, 2011; published OnlineFirst February 22, 2011.

References


Genomic Alterations of Melanoma Subtypes


Integrative Genomics Identifies Molecular Alterations that Challenge the Linear Model of Melanoma Progression

Amy E. Rose, Laura Poliseno, Jinhua Wang, et al.