Genetic evolution of T-cell resistance in the course of melanoma progression

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No potential conflicts of interest were disclosed.
Translational relevance

Metastatic melanoma is an aggressive, frequently deadly disease. In recent years immunotherapies with anti-CTLA-4 and anti-PD-1 antibodies, exploiting the capacity of CD8+ T cells to kill immunogenic tumor cells, have shown great clinical promise. However, not all patients benefit from immunotherapy and currently biomarkers predicting responses to treatment are lacking. In our study, a progressive loss in immunogenicity was observed for melanoma cells derived from three consecutive patient metastases. This culminated in complete T-cell resistance of the tumor cells due to loss of HLA class I expression. Genetic analysis revealed an early acquired chromosomal deletion and subsequent inactivating gene mutation leading to β2m deficiency and the observed HLA class I-negative phenotype. Our findings suggest that tumors can genetically evolve to avoid being recognized by the immune system. Detecting genetic alterations impacting tumor immunogenicity could be of significant value in determining which patients are likely to benefit from melanoma immunotherapy.
Abstract

**Purpose:** CD8⁺ T lymphocytes can kill autologous melanoma cells but their activity is impaired when poorly immunogenic tumor phenotypes evolve in the course of disease progression. Here, we analyzed three consecutive melanoma lesions obtained within one year of developing stage IV disease for their recognition by autologous T cells.

**Experimental Design:** One skin (Ma-Mel-48a) and two lymph node (Ma-Mel-48b, Ma-Mel-48c) metastases were analyzed for T-cell infiltration. Melanoma cell lines established from the respective lesions were characterized, determining the T cell-stimulatory capacity, expression of surface molecules involved in T-cell activation and specific genetic alterations affecting the tumor-T cell interaction.

**Results:** Metastases Ma-Mel-48a and Ma-Mel-48b, in contrast to Ma-Mel-48c, were infiltrated by T cells. The T cell-stimulatory capacity was found to be strong for Ma-Mel-48a, lower for Ma-Mel-48b and completely abrogated for Ma-Mel-48c cells. The latter proved to be HLA class I-negative due to an inactivating mutation in one allele of the beta-2-microglobulin (B2M) gene and concomitant loss of the other allele by a deletion on chromosome 15q. The same deletion was already present in Ma-Mel-48a and Ma-Mel-48b cells pointing to an early acquired genetic event predisposing to development of β2m deficiency. Notably, the same chronology of genetic alterations was also observed in a second β2m-deficient melanoma model.

**Conclusion:** Our study reveals a progressive loss in melanoma immunogenicity during the course of metastatic disease. The genetic evolvement of T-cell resistance suggests screening tumors for genetic alterations affecting immunogenicity could be clinically relevant in terms of predicting patient responses to T cell-based immunotherapy.
Introduction

CD8+ T lymphocytes (CTL) can exert potent in vivo cytotoxicity against autologous melanoma cells. This is demonstrated by the remarkable clinical responses observed in adoptive cellular therapy with autologous tumor-reactive T cells and therapy with immune-modulatory antibodies that release T cells from suppressive signals (1-5). The efficacy of these treatment regimens is based on the high intrinsic immunogenicity of melanoma cells that allows CTL to respond to multiple HLA class I-restricted tumor antigens (6). However, only a subgroup of patients receiving immunotherapy experiences clinical benefit, while others do not respond at all or after initial responses show progressive disease. In particular the coexistence of therapy-responding and non-responding metastases in individual patients suggests heterogeneous T-cell responsiveness of the different lesions (1). The underlying mechanisms therefore are most likely multi-factorial. An immunosuppressive tumor microenvironment generated by regulatory T cells, myeloid derived suppressor cells or other immune regulators can restrict T-cell effectiveness (7-9). Furthermore T-cell activity can be hampered by specific genetic alterations that affect tumor immunogenicity. Interestingly, the anti-tumor activity of the T cells is considered a driving force that selectively favors the outgrowth of low-immunogenic melanoma phenotypes. Thus, besides eliminating tumor cells, T cells shape the immunogenicity of malignant cells thereby supporting the escape of tumors from immune surveillance. This so-called T cell-mediated immunoediting of tumors, initially defined by Robert Schreiber and colleagues, has been observed in several mouse model studies (10) and evidences that this process also takes place in cancer patients have been provided. By investigating metastatic cell lines and in some cases tissue samples several genetic alterations affecting the tumor cells’ immunogenicity have been defined, including the loss of single or multiple HLA alleles and HLA haplotypes as well as the total lack of HLA class I antigen expression (11-15). So far, only a few studies followed the course of melanoma immunogenicity during disease progression due to the poor availability of tissue from...
sequential tumor metastases, corresponding cell lines and autologous peripheral blood T lymphocytes. Coulie and colleagues were the first to examine T-cell recognition of autologous tumor cells derived from two sequential metastatic melanoma lesions. They observed that T-cell responses towards melanoma cells from the second metastases were impaired as these cells expressed only one of the six HLA class I alleles (16). In 2005, Yamshchikov et al. demonstrated that recognition of tumor cells from two consecutive melanoma lesions by autologous differentiation antigen-specific CD8\(^+\) T cells was compromised by an HLA haplotype loss in tumor cells from the first metastasis and by a mixed HLA-low and HLA-negative tumor-cell phenotype in the second metastasis (14). Hence, as a basis for improving melanoma immunotherapy, more information is needed regarding the type and the sequence of genetic alterations in the tumor cells with potential impact on their recognition by cognate T cells.

Here we monitored the immunogenicity of three consecutive melanoma metastases from patient Ma-Mel-48 obtained within one year of developing stage IV disease. We observed a gradual loss in immunogenicity culminating in complete T-cell resistance of the tumor cells caused by an irreversible HLA class I-negative phenotype. This originated from two sequential genetic alterations, a deletion on chromosome 15q where the \(B2M\) gene maps and an inactivating \(B2M\) gene mutation. Notably, tumor cells from all metastases of patient Ma-Mel-48 carried the same deletion on chromosome 15q, identifying this alteration as an early predisposing genetic event in the development of \(\beta2\)m deficiency. The same chronology of genetic alterations was also observed in a second patient model of \(\beta2\)m-deficient melanoma.
Materials and Methods

Patient samples

Patient Ma-Mel-48 presented with melanoma stage II in December 2000 at the age of 85 years. Progression to stage III and IV was diagnosed in October 2001 and June 2002, respectively. The patient was treated with Temozolomide at the beginning of 2003 but never received immunotherapy. In September 2003 the patient died. Patient Ma-Mel-100 presented with melanoma stage II in January 1998 at the age of 72 years. Progression to stage III was diagnosed in September 2002. From February 1998 until September 2002 the patient received adjuvant interferon (IFN)-α therapy. In September 2005 the patient died.

Samples including tumor tissues and peripheral blood mononuclear cells (PBMC) were collected after approval by the institutional review board and patient informed written consent. Tissues were mechanically divided for cryopreservation and generation of the corresponding cell lines. Small tissue pieces were distributed in cell culture dishes and outgrowing cells were split for the first time at 90% cell confluence. Melanoma cell lines were cultured in RPMI1640 medium supplemented with glutamine, 10% FCS and penicillin/streptomycin. Cells were cultured at 37°C in a 5% CO₂ atmosphere. Cell lines were authenticated by genetic profiling at the Institute for Forensic Medicine (University Hospital Essen) using the AmpFLSTR-Profiler Plus kit (Applied Biosystems) and routinely tested every 6 months.
Antibodies

The following murine monoclonal antibodies (mAb) were used for immunohistochemistry: W6/32 to detect HLA class I antigens (Dianova); bbm.1 to stain for β2m (kindly provided by G. Moldenhauer, German Cancer Research Center, Heidelberg, Germany); anti-HMB-45 (Dako) to detect melanoma cells; anti-CD3 (BD Pharmingen) to stain for T cells.

For flow cytometry the mouse mAb were as follows: anti-HLA-ABC-APC (eBiosciences), anti-CD54-PE and anti-HLA-DR-PECy7 (Beckmann Coulter), anti-PD-L1-PE and anti-PD-L2 (Biolegend), anti-B7-H3 (R&D Systems), anti-B7-H4 (eBiosciences); L243 was used for detection of HLA-DR molecules (17) and HC10 for labelling of β2m-free HLA heavy chains (18, 19).

The following antibodies were used for Western blot analysis: mouse anti-Melan-A/MART-1 (Zytomed), mouse anti-tyrosinase and anti-MITF (Santa Cruz), rabbit anti-DCT/TRP2 (kindly provided by V. Hearing, National Cancer Institute, National Institutes of Health, Bethesda); mouse anti-STAT1, rabbit anti-phospho(p)STAT1, rabbit anti-JAK1, rabbit anti-GAPDH (Cell Signaling); rabbit anti-IRF1 and mouse anti-β2m (Santa Cruz); rabbit anti-β2m (Sigma); mouse anti-TAP1 (TO-1) and mouse anti-tapasin (TO-3) (20); mouse mAb HC10 (18, 19).

Mixed lymphocyte-tumor cell culture (MLTC)

Anti-CD8-Microbeads (Miltenyi Biotec) were used for positive selection of CD8+ T cells from PBMC. Selected T cells were cocultured in 24-well plates at 10⁶ cells/well with irradiated (100 Gray) autologous tumor cells at 10⁵ cells/well in 2 mL AIM-V medium (GIBCO-BRL) supplemented with 10% human AB serum (complete medium). Interleukin-2 (IL-2) was added on day 3 at 250 IU/mL (Chiron Corporation). Lymphocytes (10⁶ cells/well) were restimulated weekly with 10⁵ irradiated tumor cells in IL-2 supplemented complete medium.
Intracellular cytokine staining

For detection of intracellular IFN-γ and TNF-α, lymphocytes were stimulated for 4 h with the indicated tumor cells (effector to target ratio of 1:1 or 1:2) in AIM-V complete medium containing 10 µg/mL Brefeldin A (Sigma-Aldrich). Then cells were stained with anti-human CD3-PE/Cy7 and CD8-APC-Alexafluor700 antibodies (Beckman Coulter) followed by fixation and permeabilization using the Fixation/Permeabilization Concentrate and Diluent kit (eBioscience) and addition of an anti-IFN-γ-FITC antibody (Beckman Coulter) or anti-TNF-α-Pacific Blue antibody (Biolegend). Cells were analysed in a Gallios flow cytometer and the Kaluza software was used for data analysis (Beckman Coulter). Where indicated, antibody W6/32 (50 µg/ml; purified from hybridoma supernatant, kindly provided by M. Fatho, Mainz) and anti-PD-L1 (10 µg/ml; Biolegend) were added in order to block the TCR/HLA class I and PD-1/PD-L1 interactions, respectively. Mouse monoclonal IgG1 (mIgG) (R&D Systems) was used as control antibody.

SNP array analysis

Genomic DNA was isolated from melanoma cell lines of patients Ma-Ma-48 and Ma-Mel-100 and from non-fixed peripheral blood cells (available only for patient Ma-Mel-48) using the QIAamp DNA Mini Kit (Qiagen, Hilden, Germany) according to the manufacturer’s instructions. SNP arrays were performed using CytoScan® HD Array from Affymetrix. Hybridization was done according to manufacturer’s protocol and data analysis performed applying the program Chromosome Analysis Suite® from Affymetrix. SNP array data files are accessible at the NCBI GEO database (accession number: GSE60218).

In order to infer the phylogenetic relationship of Ma-Mel-48 cells, genotype calls were extracted from the CytoScan® HD Array data using the linux shell and R (21). As an out-group genotype calls from the autologous blood sample (166) were used. We identified 111029 variable sites in which the genotype calls of the cell lines tested and the blood sample
differed from each other, 64023 of those were parsimony informative. We then used the maximum parsimony criterion, minimizing the total number of evolutionary steps required to explain the relationship of the tested samples. We ran maximum parsimony in MEGA (22) using 500 bootstrap replications and complete deletion of missing data, as well as subtree-pruning-regrafting as a search method.

Additional method information is provided as supplemental material.
Results

Consecutive metastases from patient Ma-Mel-48 show heterogeneous T-cell infiltration and HLA class I antigen expression

To determine the development of melanoma immunogenicity during disease progression we collected tumor tissues from three consecutive metastases of patient Ma-Mel-48. The first Ma-Mel-48a, a skin metastasis, was excised in July 2002, a few weeks after diagnosis of stage IV disease. The second and the third lesions Ma-Mel-48b and Ma-Mel-48c, lymph node metastases, were surgically removed in January 2003 and July 2003, respectively (Fig. 1a).

Serial tissue sections of the three metastases were first analyzed by immunohistochemistry for their infiltration by CD3$^+$ T lymphocytes. T cells accumulated in the periphery and center of metastases Ma-Mel-48a and Ma-Mel-48b pointing to a close tumor-T cell interaction (Fig. 1b and Supplementary Fig. S1). In contrast, no T cell infiltrate was detected in metastasis Ma-Mel-48c (Fig. 1b). The presence of T cells correlated with the expression of HLA class I molecules, stained for with the antibody W6/32 binding a structural epitope formed by complexes of the β2m light chain and the HLA heavy chains. HMB-45-positive melanoma cells expressed HLA class I molecules in metastasis Ma-Mel-48a but not in Ma-Mel-48c (Fig. 1b).

Next, cell lines established from the respective metastatic lesions were analyzed for their HLA class I surface expression by flow cytometry. Ma-Mel-48a as well as Ma-Mel-48b cells were MHC class I-positive while Ma-Mel-48c cells proved to be negative (Fig. 1c). Thus, the data generated by flow cytometry were in agreement with the results obtained by immunohistochemistry.

Progressive loss in T-cell recognition of the different melanoma cell lines
On basis of the HLA class I expression pattern we analyzed the T cell-stimulatory capacity of Ma-Mel-48a and Ma-Mel-48b cells. MLTC were set up, consisting of purified autologous peripheral blood CD8\(^+\) T cells and Ma-Mel-48a or Ma-Mel-48b tumor cells. CD8\(^+\) T cells were stimulated in weekly intervals with the indicated irradiated tumor cells to increase the frequency of tumor-reactive T lymphocytes. After two stimulations CD8\(^+\) T cells were harvested and analyzed for their responsiveness towards the tumor cells. As shown in Fig. 2a, approximately 9% of CD8\(^+\) T cells from MLTC with Ma-Mel-48a cells (MLTC-48a) responded to Ma-Mel-48a cells with IFN-\(\gamma\) production, while only around 2% of the CD8\(^+\) T cells reacted against Ma-Mel-48b cells. Interestingly, CD8\(^+\) T cells harvested from MLTC-48b showed limited responsiveness to Ma-Mel-48b as well as Ma-Mel-48a cells (Fig. 2b). Approximately 2% of the T cells recognized Ma-Mel-48b cells and comparable reactivity towards Ma-Mel-48a cells was observed. These results pointed to a lower T cell-stimulatory capacity of Ma-Mel-48b cells as compared to Ma-Mel-48a cells. In either case, production of CD8\(^+\) T-cell cytokines proved to be HLA class I-dependent (Fig. 2c), thus no T-cell reactivity towards the HLA class I-negative Ma-Mel-48c cells was observed (Fig. 2a-b).

In contrast to CD8\(^+\) T cells, autologous CD4\(^+\) T cells did not react towards any of the Ma-Mel-48 cell lines that lacked HLA class II expression even in the presence of IFN-\(\gamma\), due to epigenetic silencing of the transcriptional regulator CIITA (Supplementary Fig. S2 and data not shown).

Low immunogenic phenotype of Ma-Mel-48b cells is not due to altered expression of HLA alleles or immunomodulatory B7 molecules

To define the molecular mechanisms underlying differential T-cell recognition we first analyzed Ma-Mel-48a and Ma-Mel-48b cells for potential alterations in HLA allele expression. HLA genotype analysis demonstrated that the HLA-A*0101, -B*0801, -C*0701 homozygous haplotype of autologous peripheral blood cells was conserved in the cell lines,
both expressing considerable amounts of HLA-A, HLA-B and HLA-C specific mRNA (Fig. 3a). Data generated by qRT-PCR pointed to elevated HLA-B mRNA levels in Ma-Mel-48b cells as compared to Ma-Mel-48a cells. Accordingly, staining of Ma-Mel-48b cells for surface expression of HLA class I antigens and β2m-free HLA heavy chains was more intense compared to Ma-Mel-48a cells (Fig. 1c, 3a). Not expecting this shift in HLA class I allele expression to be responsible for the lower immunogenic phenotype of Ma-Mel-48b cells, we studied the surface expression of additional molecules involved in tumor-T cell interaction. As shown in Fig. 3b, both cell lines expressed similar levels of the immunomodulatory B7 molecules PD-L1 (B7-H1), PD-L2 (B7-DC) and B7-H3, whereas no expression of B7-H4 was noted in either case. Though inhibitory PD-L1 was clearly detectable on Ma-Mel-48a and Ma-Mel-48b cells its blockade did not enhance T-cell activities in our in vitro settings (Supplementary Fig. S3). Both melanoma cell lines also expressed similar levels of CD54 (ICAM-1), of importance for T-cell adhesion to the tumor cells (Fig. 3c).

While the analysis of specific surface molecules did not reveal significant differences between Ma-Mel-48a and Ma-Mel-48b cells we postulated that variations in antigen expression levels could be responsible for the differential immunogenicity. Although the specificity of tumor-reactive T cells could not be studied due to limited sample material, we investigated the expression of melanoma differentiation antigens (MDA) in the tumor cells. As shown in Fig. 3d, Melan-A, tyrosinase and TRP2/DCT as well as their transcriptional regulator MITF were decreased in Ma-Mel-48b as compared to Ma-Mel-48a cells, suggesting MDA downregulation in Ma-Mel-48b cells could contribute to their lower T cell-stimulatory capacity. Interestingly, Ma-Mel-48b dedifferentiation was not associated with a slow proliferative phenotype, as frequently observed for MITF-low melanoma cells (Supplementary Fig. S4) (23).

The HLA class I-negative phenotype of Ma-Mel-48c cells is caused by β2m deficiency
To determine whether the HLA class I-negative phenotype of Ma-Mel-48c cells was reversible cells were treated with type I and type II IFN. As shown in Fig. 4a, surface levels of HLA class I antigens were enhanced on Ma-Mel-48a and Ma-Mel-48b cells in response to IFN treatment. However, HLA class I antigen expression on Ma-Mel-48c cells could not be restored (Fig. 4a), although basic expression of interferon pathway components in Ma-Mel-48c cells was similar to the other cell lines and also the upregulation of STAT1, pSTAT1 and IRF1 in response to interferon treatment was comparable (Fig. 4b, Supplementary Fig. S5a).

Analysis of several components of the antigen processing and presentation machinery (APM) by Western blot revealed expression of HLA heavy chains, TAP1 and tapasin in all cell lines while β2m protein was expressed in Ma-Mel-48a and Ma-Mel-48b but not in Ma-Mel-48c cells, even when cells were treated with IFN (Fig. 4b). Also mRNA expression of specific APM components was comparable between the three melanoma cell lines and was detectable in similar amounts also in melanocytes, but only Ma-Mel-48c cells lacked B2M mRNA (Fig. 4c, Supplementary Fig. S5b). These findings suggested the HLA class I-negative phenotype of Ma-Mel-48c cells was caused by the lack of B2M gene expression. Although B2M mRNA was not detectable by qRT-PCR, we obtained a shortened B2M-specific product by RT-PCR only for Ma-Mel-48c cells with primers amplifying the coding region from the start to the stop codon (Fig. 5a). Sequence analysis of this PCR product identified a 60 bp deletion starting in codon 96 (Exon II) of the B2M gene, leading to a C-terminally altered gene product (Fig. 5b). Notably, this B2M deletion variant could not be detected by qRT-PCR (Fig. 4c), as the assay primers were located in the B2M Exon II-III boundary. To confirm that the loss in HLA class I antigen expression was solely due to β2m deficiency, Ma-Mel-48c cells were transfected with a B2M expression plasmid. As shown in Fig. 5c, induction of HLA class I antigen expression was detectable on a population of transiently B2M-transfected cells. In agreement with the results obtained for the cell line, β2m was not detected in tumor cells in the corresponding metastatic lesion while it was expressed by normal cells (Fig. 5d).
**B2M allele loss is an early predisposing genetic event in the development of β2m deficiency**

Previously, we reported that β2m deficiency is caused by the coincidence of a B2M gene mutation and allelic B2M loss. The B2M gene maps to chromosome 15q21.1 and can be lost as a result of chromosome 15q aberrations (12, 24). To define whether this was also the case in Ma-Mel-48c cells, SNP array analyses were performed on DNA obtained from the various tumor cell lines and autologous peripheral blood cells as a constitutive, normal control. Indeed, a partial deletion on chromosome 15q was observed in Ma-Mel-48c cells, encompassing the region 15q13.3 to 15q21.2 (Fig. 5e). Interestingly, the same portion of chromosome 15q was also found deleted in Ma-Mel-48a and Ma-Mel-48b cells.

From the genome-wide SNP genotype data we therefore inferred the phylogenetic relationship of the melanoma cell lines and the corresponding autologous blood sample. A phylogenetic tree based on maximum parsimony showed that Ma-Mel-48b and Ma-Mel-48c cells were more closely related to each other than to Ma-Mel-48a cells indicating that a lineage leading to Ma-Mel-48a evolved first. The grouping was present in all 500 bootstrap replicates of the analysis (Fig. 5f). Overall these results demonstrated that the allelic loss of B2M occurred early in the course of disease and that a subsequent 60 bp deletion in the remaining B2M allele resulted in β2m deficiency in Ma-Mel-48c cells.

**β2m-deficient melanoma cells from patient Ma-Mel-100 show the same chronology of genetic alterations**

In addition to patient Ma-Mel-48, we observed loss of HLA class I antigen expression on melanoma cells from patient Ma-Mel-100 (Fig. 6a). Tumor cells from the two regional lymph node metastases Ma-Mel-100a and Ma-Mel-100b excised in April 2004 and May 2005, respectively, could not be stained with mAb W6/32 (Fig. 6b, Supplementary Fig. S6a).
Consistent with the lack of HLA class I expression, T cells did not infiltrate the tumor lesion but were localized in the periphery (Fig. 6b). Also cell lines established from the metastases showed an HLA class I-negative phenotype, which remained stable in the presence of type I and II interferons (Fig. 6c, data not shown). But HLA class I surface expression on Ma-Mel-100a and Ma-Mel-100b cells could be induced upon transfection of a \textit{B2M} expression plasmid (Fig. 6c). Both cell lines also expressed similar amounts of \textit{B2M}-specific mRNA (Fig. 6d) however, only Ma-Mel-100a cells lacked \(\beta\)2m expression \textit{in situ} and \textit{in vitro} (Fig. 6b, e), while the protein was detected in lysates from Ma-Mel-100b cells (Fig. 6e).

Screening of the cells by SNP array for the underlying molecular alterations revealed deletion of the same region on chromosome 15q in Ma-Mel-100a and Ma-Mel-100b cells, ranging from 15q12 to 15q22.2 (Fig. 6f). Thus, the shared loss of one \textit{B2M} allele was the initial genetic alteration in the development of \(\beta\)2m deficiency. Additional alterations, acquired subsequently, were different in both cell lines. Sequence analysis of the coding region revealed a 12 bp deletion in Exon II of the \textit{B2M} gene in Ma-Mel-100b cells. In contrast, only wild type \textit{B2M} mRNA was detected in Ma-Mel-100a cells, pointing to an unknown post-transcriptional defect in \textit{B2M} expression (Fig. 6f, Supplementary Fig. S6b).
Discussion

During disease progression melanoma cells acquire genetic alterations that can affect their recognition by cytotoxic T cells. While some mutations increase the T cell sensitivity of melanoma cells (25-27) others decrease the tumor cell’s immunogenicity and interfere with the effectiveness of immunotherapies. In terms of predicting responses to T cell-based immunotherapy it is important to understand the type and the sequence of genetic alterations that hamper the recognition of melanoma cells by cognate T cells.

Here we studied the T cell-stimulatory capacity of melanoma cells from three consecutive metastases of patient Ma-Mel-48 obtained at different times within one year of progressive stage IV disease. While melanoma cells from skin metastasis Ma-Mel-48a strongly stimulated autologous CD8+ T cells, tumor cells from the lymph node metastasis Ma-Mel-48b cells, excised half a year later, did not. The lower T cell-stimulatory capacity of Ma-Mel-48b cells could not be explained by an altered expression of immunomodulatory B7 molecules or the adhesion molecule CD54. In addition, there was no expression of HLA-DR molecules, described as ligands of the regulatory receptor LAG3 (28-31). Furthermore, HLA allele alterations, known to protect melanoma cells from certain T-cell specificities, were not detected in Ma-Mel-48b cells (13, 15, 32). We therefore assume the lower immunogenicity of Ma-Mel-48b cells was more likely due to differences in the antigen expression pattern. Specific T-cell antigens could be down-regulated in Ma-Mel-48b in comparison to Ma-Mel-48a cells, as observed for the MDA (33). Furthermore, alterations in proteasome subunits and peptidases involved in the processing and presentation of tumor antigens could have contributed to the differential immunogenicity of the tumor cells (34-36).

The gradual decrease of melanoma immunogenicity in patient Ma-Mel-48 culminated in the complete T-cell resistance of Ma-Mel-48c cells, as these cells acquired an irreversible HLA class I-negative phenotype. This was caused by a lack in β2m expression due to an inactivating mutation in one allele of the B2M gene and concomitant loss of the second allele.
We previously reported that β2m-deficient melanoma cells are characterized by deletions on chromosome 15q including the \textit{B2M} gene mapping to 15q21.1 (24). Notably, loss of chromosome 15q material in melanoma is not a very rare event (37, 38). Of 70 metastatic melanoma samples analyzed 16% were positive for loss of heterozygosity in chromosome region 15q21-22 (39). In the near future, exome studies will provide more detailed information about the frequency of \textit{B2M} gene mutations in melanoma metastases, which we expect to be lower than \textit{B2M} allelic losses.

By SNP array analysis on DNA from Ma-Mel-48c cells we could clearly narrow down the deletion on chromosome 15q including the \textit{B2M} gene. Notably, exactly the same deletion on chromosome 15q was already present in Ma-Mel-48a and Ma-Mel-48b cells. A phylogenetic tree inferred from SNP array data demonstrated that in patient Ma-Mel-48 a lineage leading to metastasis Ma-Mel-48a cells evolved first, followed by the lineages leading to Ma-Mel-48b and Ma-Mel-48c. This branching order is consistent with the timing of the excision of the metastases. The localized deletion on chromosome 15q, including one \textit{B2M} allele, detected in all metastatic cell lines is thus an early event on the lineage leading to all three studied melanoma metastases. To our knowledge this is the first study demonstrating that allelic \textit{B2M} loss is the initial genetic alteration in the development of β2m deficiency. The same chronology of genetic alterations was also observed in tumor cells of patient Ma-Mel-100. Here two metastases shared a common deletion on chromosome 15q associated with loss of a \textit{B2M} allele. Remarkably, both metastases were found to be β2m-deficient due to different subsequent alterations. A specific \textit{B2M} gene mutation was identified in Ma-Mel-100b cells, whereas Ma-Mel-100a cells expressed significant amounts of only wild type \textit{B2M} mRNA. We assume that an unknown post-transcriptional mechanism blocks \textit{B2M} mRNA translation in Ma-Mel-100a cells that requires further investigations.

While our data suggest deletion of a \textit{B2M} allele can be an early event in the course of melanoma progression, most studies detected β2m-deficient melanoma cells in late stage
disease (12, 14). Interestingly, Del Campo et al. recently found nests of β2m-negative tumor cells to be present in a very early lesion from a melanoma patient, who later developed a completely HLA class I-negative metastasis (40). While the first β2m-negative tumor cells were detectable prior to immunotherapy, the completely HLA class I-negative lesion occurred under dendritic cell vaccination leading to the assumption that vaccine-induced T-cell activity enriched β2m-deficient tumor cells. This was proposed also by previous studies that detected β2m loss in several recurrent metastases from patients receiving immunotherapy, such as patient Ma-Mel-100 who was treated with IFN-α (13, 41). Since patient Ma-Mel-48 did not receive immunotherapy this suggests that spontaneous anti-tumor T-cell responses led to the outgrowth of the HLA class I-negative melanoma immunophenotype. Besides, T cell-independent mechanisms should also be taken into consideration. Recently, Garrido and colleagues demonstrated that HLA class I-negative melanoma cells showed enhanced proliferation, migration and invasion in comparison to their HLA class I-positive counterparts, pointing to HLA class I molecules as tumor suppressors (42). Irrespective of the driving force, HLA class I alterations, in particular HLA class I loss, impede T-cell recognition of melanoma and will negatively influence all T cell-based immunotherapies.

The infiltration of melanoma metastases by T cells is currently discussed as a biomarker that predicts responsiveness to immunotherapy, in that patients with T cell-inflamed metastases more likely respond to treatment (43-45). Different mechanisms determine the migration of T cells into a metastatic lesion, such as the release of chemokines by tumor cells (45, 46). Furthermore, it has been described that the intensity of the T-cell infiltrate correlates with the level of HLA class I antigens expressed in the tumor (47). In accordance, CD3+ T cells were not detected in the center of metastasis Ma-Mel-48c or Ma-Mel-100a. T cells were rather located at the periphery, an observation also noted in other HLA class I-deficient melanoma lesions (unpublished data). This suggests that within the group of melanoma patients with non T cell-inflamed tumors a subgroup might exhibit HLA abnormalities. Thus we propose that
biopsies from melanoma patients should be screened for genetic HLA alterations, including losses on chromosome 15q21.1, as this could prove valuable for immunotherapeutic treatment decisions.
Literature


33. Yee C, Thompson JA, Byrd D, Riddell SR, Roche P, Celis E, et al. Adoptive T cell therapy using antigen-specific CD8+ T cell clones for the treatment of patients with...


Figure Legends

**Fig 1: Heterogeneous T-cell infiltration and HLA class I antigen expression in different metastases from patient Ma-Mel-48**

A, localization and excision date of three sequential metastases from patient Ma-Mel-48. B, serial sections from cryopreserved Ma-Mel-48a and -48c metastatic lesions were analysed for the expression of HMB-45 (melanoma cells), CD3 (T cells) and HLA class I antigens by immunohistochemistry. Red staining indicates positive cells. C, flow cytometric analysis of total HLA class I antigen expression on melanoma cell lines established from the different metastases of patient Ma-Mel-48. Cells were stained with mAb W6/32. Histograms from one representative of three independent experiments are shown. Numbers indicate mean fluorescence intensity (MFI).

**Fig 2: Tumor cells from sequential lesions differ in their T cell-stimulatory capacity**

A-C, analysis of the T cell-stimulatory capacity of the different Ma-Mel-48 tumor cell lines. In autologous MLTC isolated CD8^+ T cells were stimulated twice with Ma-Mel-48a (MLTC-48a; A, C) or Ma-Mel-48b (MLTC-48b; B, C) cells and subsequently analysed for their response towards different target cells by intracellular staining for IFN-γ or TNF-α. A left, first dot plot: spontaneous IFN-γ production by CD8^+ T cells from MLTC-48a; second to fourth dot plot: production of IFN-γ in response to the different target cells indicated above. B left, first dot plot: spontaneous IFN-γ production by CD8^+ T cells from MLTC-48b; second to fourth dot plot: production of IFN-γ in response to the different target cells indicated above. Results from one representative of two independent experiments are presented, numbers in dot plots indicate %IFN-γ^+CD8^+ T cells within CD8^+CD3^+ T cells. C, to demonstrate HLA class I-dependent production of T-cell cytokines, tumor cells were incubated with blocking mAb W6/32 before addition of T cells. Control cells were incubated with mouse IgG1.
(mIgG). The mean %TNF-α+CD8+ T cells within CD8+CD3+ T cells of n=2 or n=3 independent experiments is presented.

**Fig. 3: Conserved HLA phenotype but altered antigen expression in Ma-Mel-48b cells**

A left, mRNA levels of *HLA-A, HLA-B, HLA-C* in Ma-Mel-48a and Ma-Mel-48b cells were quantified by qRT-PCR and normalized to endogenous *GAPDH* mRNA. Expression levels, given as means (+ SEM) of three independent experiments, are depicted relative to the expression in Ma-Mel-48a cells. A right, flow cytometric analysis of β2m-free HLA heavy chain expression on Ma-Mel-48a and Ma-Mel-48b cells with mAb HC10. B, analysis of PD-L1 (B7-H1), PD-L2 (B7-DC), B7-H3, B7-H4 and CD54 surface expression on melanoma cell lines Ma-Mel-48a and Ma-Mel-48b by flow cytometry. Histograms from one representative of three independent experiments are shown. C, Expression of MDA (Melan-A, tyrosinase, TRP2/DCT) and its transcriptional regulator MITF in Ma-Mel-48a and Ma-Mel-48b cells was determined by Western blot. GAPDH served as a loading control. Data from one representative of three independent experiments are presented.

**Fig 4: Interferon treatment of Ma-Mel-48c cells does not restore HLA class I antigen surface expression**

The different Ma-Mel-48 cell lines were treated with IFN-α (1000 U/ml) or IFN-γ (500 U/ml) for 48 h, controls were left untreated. A, cells were stained with mAb W6/32 for detection of HLA class I antigen expression. Data from one representative of three independent experiments are depicted, numbers indicate MFI values. B, cell lysates were analyzed by Western blot for the protein levels of interferon pathway components (STAT1, pSTAT1, IRF1) or APM components (HLA heavy chains, tapasin, TAP1, β2m). GAPDH served as loading control. One representative of three independent experiments is depicted. C, mRNA levels of different APM components were quantified by qRT-PCR and normalized to
endogenous *GAPDH* mRNA. Expression levels, given as means (+ SEM) of three independent experiments, are depicted relative to the expression in Ma-Mel-48a cells.

**Fig 5: β2m deficiency of Ma-Mel-48c cells is caused by B2M gene mutation subsequent to B2M allele loss**

A, semi-quantitative analysis of *B2M* mRNA expression by RT-PCR. Ma-Mel-48c cells express a shortened *B2M* mRNA, a PCR product in the size of wild type (wt) mRNA was only detectable in Ma-Mel-48a and Ma-Mel-48b cells. B, comparison of the C-terminal amino acid sequence (aa) of mutant β2m, as expressed in Ma-Mel-48c cells, and wt β2m. Capital letters indicate the aa sequence. The first 95 aa are identical in wt and mutant proteins. Stars indicate stop codons; C, Ma-Mel-48c cells were transfected with a *B2M* expression plasmid. Expression of HLA class I antigens on transient transfectants was determined by flow cytometry. D, analysis of β2m expression in metastatic lesion Ma-Mel-48c by immunohistochemistry. Tumor cells were negative for β2m and did not stain red. E, SNP results given as allelic distribution of chromosome 15q are shown for DNA obtained from autologous peripheral blood mononuclear cells and melanoma cells. All Ma-Mel-48 cell lines show loss of one chromosomal allele in the region 15q13.3 - 15q21.2 (Chr.15: 33,045,756-50,579,508; hg19). The location of *B2M* at 15q21.1 (Chr.15: 45,003,675-45,011,075) is shown by the dashed line. F, maximum parsimony tree showing the phylogenetic relationship of the melanoma cell lines and the blood sample (166) used as outgroup. Within the patient, a melanoma lineage leading to the studied cell lines evolved and was genetically divergent from the blood sample. A *B2M* loss evolved on this lineage. Then, a lineage leading to Ma-Mel-48a diverged from the melanoma ancestor and cumulated genotypic differences specific to this lineage. Later the lineages of Ma-Mel-48b and Ma-Mel-48c diverged and cumulated their specific genotypic differences. 100 percent of the bootstrap replicates showed this grouping.
Fig 6: B2M allele loss is also the initial genetic alteration in β2m-deficient melanoma cells from patient Ma-Mel-100

A, location and excision date of two sequential metastases from patient Ma-Mel-100. B, serial sections from cryopreserved metastasis Ma-Mel-100a were analysed for the expression of HMB-45 (melanoma cells), CD3 (T cells), HLA class I antigens and β2m by immunohistochemistry. Red staining indicates positive cells. C, flow cytometric analysis of total HLA class I antigens on cell lines established from metastases Ma-Mel-100a and Ma-Mel-100b before and after transient transfection with a B2M expression plasmid. Cells were stained with mAb W6/32. Representative histograms from one of three independent experiments are shown. D, B2M mRNA levels were quantified by qRT-PCR and normalized to endogenous GAPDH mRNA. Expression levels, given as means (+ SEM) of three independent experiments, are depicted relative to the expression in Ma-Mel-100b cells. E, β2m protein levels in cell lysates were determined by Western blot. GAPDH served as loading control. Representative results from one of three independent experiments are depicted. F, SNP array results showing the allelic distribution of chromosome 15q from DNA obtained from Ma-Mel-100a and Ma-Mel-100b cells. One chromosomal allele in the region 15q12 to 15q22.2 (Chr.15: 26,840,997-61,436,881; hg19) is lost in both Ma-Mel-100 cell lines. The location of B2M at 15q21.1 is shown by the dashed line. A subsequent B2M gene mutation was only detected in Ma-Mel-100b cells, identified as 12 bp deletion in Exon II.
A

<table>
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<th>Tumor</th>
<th>Metastasis</th>
<th>Excision</th>
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<tbody>
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<tr>
<td>Ma-Mel-48c</td>
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B

Ma-Mel-48a

\[
\text{HMB-45} \quad \text{CD3} \quad \text{HLA class I}
\]

![Images of HMB-45, CD3, and HLA class I for Ma-Mel-48a with magnifications 5x and 10x.]

Ma-Mel-48c

\[
\text{HMB-45} \quad \text{CD3} \quad \text{HLA class I}
\]

![Images of HMB-45, CD3, and HLA class I for Ma-Mel-48c with magnifications 5x and 10x.]

C

Ma-Mel-48a

\[
\text{Events: 24.29}
\]

Ma-Mel-48b

\[
\text{Events: 31.66}
\]

Ma-Mel-48c

\[
\text{Events: 0.82}
\]

![Graphs showing HLA class I events for Ma-Mel-48a, Ma-Mel-48b, and Ma-Mel-48c.]

Fig. 1
A

<table>
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B

C

Fig. 2

Author Manuscript Published OnlineFirst on October 7, 2014; DOI: 10.1158/1078-0432.CCR-14-0567

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Fig. 3

A

**HLA-A**

**HLA-B**

**HLA-C**

relative mRNA levels

Ma-Mel-48a Ma-Mel-48b

Ma-Mel-48a Ma-Mel-48b

β2m-free HLA heavy chain

B

**Ma-Mel-48a**

**Ma-Mel-48b**

Events

PD-L1 (B7-H1)

Ma-Mel-48a Ma-Mel-48b

Ma-Mel-48a Ma-Mel-48b

PD-L2 (B7-DC)

Ma-Mel-48a Ma-Mel-48b

Ma-Mel-48a Ma-Mel-48b

B7-H3

B7-H4

C

**Ma-Mel-48a**

**Ma-Mel-48b**

Events

CD54

Ma-Mel-48a Ma-Mel-48b

D

**Ma-Mel-48a**

**Ma-Mel-48b**

Melan-A

DCT/TRP2

tyrosinase

MITF

GAPDH

Fig. 3
Fig. 4
Fig. 5

A

B2M

GAPDH

B

\[ \beta_2m\text{ wt } KDEYACR\text{VNHVTL...}^* \]

\[ \beta_2m\text{ mutant } KVSFS^* \]

(Ma-Mel-48c)

C

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</tr>
<tr>
<td>10^4</td>
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<tr>
<td>HLA class I</td>
</tr>
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</table>

D

20x

E

Germline

Ma-Mel-48a

Ma-Mel-48b

Ma-Mel-48c

Chr. 15q

B2M

F

Melanoma lineage

B2M allele loss

100

48c LN metastasis

48b LN metastasis

49a Skin metastasis

165 Blood

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<table>
<thead>
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<th>Metastasis</th>
<th>Excision</th>
</tr>
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<tbody>
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<tr>
<td>Ma-Mel-100b</td>
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**B**

- HMB-45
- CD3
- HLA class I
- β2m

**C**

- Ma-Mel-100a + B2M
- Ma-Mel-100b + B2M

**D**

- relative mRNA levels

**E**

- Ma-Mel-100a
- Ma-Mel-100b
- β2m
- GAPDH

**F**

- allelic loss
- subsequent alteration
- unknown mechanism
- 12 bp deletion in Exon II

**Chr. 15q**

- B2M

Fig. 6
Genetic evolution of T-cell resistance in the course of melanoma progression

Antje Sucker, Fang Zhao, Birgit Real, et al.

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