Association of IFN-γ Signal Transduction Defects with Impaired HLA Class I Antigen Processing in Melanoma Cell Lines

Annedore Respa1, Juergen Bukur1, Soldano Ferrone3, Graham Pawelec2, Yingdong Zhao4, Ena Wang3, Francesco M. Marincola3, and Barbara Seliger1

Abstract

Purpose: Abnormalities in the constitutive and IFN-γ-inducible HLA class I surface antigen expression of tumor cells is often associated with an impaired expression of components of the antigen processing machinery (APM). Hence, we analyzed whether there exists a link between the IFN-γ signaling pathway, constitutive HLA class I APM component expression, and IFN-γ resistance.

Experimental Design: The basal and IFN-γ-inducible expression profiles of HLA class I APM and IFN-γ signal transduction cascade components were assessed in melanoma cells by real-time PCR (RT-PCR). Western blot analysis and/or flow cytometry, the integrity of the Janus activated kinase (JAK) 2 locus by comparative genomic hybridization. JAK2 was transiently overexpressed in JAK2- cells. The effect of IFN-γ on the cell growth was assessed by XTT [2,3-bis(2-methoxy-4-nitro-5-sulfophenyl)-H-tetrazolium-5-carboxanilide inner salt] assay.

Results: The analysis of 8 melanoma cell lines linked the IFN-γ unresponsiveness of Colo 857 cells determined by lack of inducibility of HLA class I surface expression on IFN-γ treatment to a deletion of JAK2 on chromosome 9, whereas other IFN-γ signaling pathway components were not affected. In addition, the constitutive HLA class I APM component expression levels were significantly reduced in JAK2- cells. Furthermore, JAK2-deficient cells were also resistant to the antiproliferative effect of IFN-γ. Transfection of wild-type JAK2 into JAK2- Colo 857 not only increased the basal APM expression but also restored their IFN-γ sensitivity.

Conclusions: Impaired JAK2 expression in melanoma cells leads to reduced basal expression of MHC class I APM components and impairs their IFN-γ inducibility, suggesting that malfunctional IFN-γ signaling might cause HLA class I abnormalities. Clin Cancer Res; 17(9): 2668–78. ©2011 AACR.
Translational Relevance

Malignant transformation of cells is frequently associated with defects in HLA class I antigen processing machinery (APM), which plays a crucial role in the recognition of tumor cells by host immune system. The potential clinical relevance of these defects has stimulated interest in the molecular characterization of the underlying mechanisms, as this information may suggest targeted therapies to restore APM function. Multiple escape mechanisms have been identified and characterized. However, little information is available about defects in the IFN-γ signal transduction pathway in malignant cells despite the susceptibility of several APM components to modulation by IFN-γ. This study has identified for the first time loss or downregulation of JAK2 as a mechanism underlying HLA class I antigen downregulation in melanoma cells and lack of inducibility by IFN-γ. Furthermore, this information contributes to the design of strategies both to identify the mechanisms underlying defects in HLA class I antigen expression in malignant cells and to correct these defects.

(IFN-γ, JAK2, and STAT1 phosphorylation upon incubation with cell line has been found to be associated with the lack of upregulation of TAP in one renal cell carcinoma (RCC) resistance have been investigated only to a limited extent, as a result, HLA class I and class II antigens are not upregulated when cells are exposed to IFN-γ (11, 12). This abnormality is likely to have a negative impact on the interactions of tumor cells with host immune system and provide them with an escape mechanism. The molecular mechanisms causing IFN-γ resistance have been investigated only to a limited extent, although this information might have an important impact on the development of targeted therapies. So far, IFN-γ-responsive genes have been shown to be frequently downregulated in tumor cells because of impaired IRF1 expression as well as defective transcriptional and posttranscriptional regulation of components of the IFN-γ signal transduction cascade. In addition, to the best of our knowledge, loss of the IFN-γ-mediated upregulation of TAP in one renal cell carcinoma (RCC) cell line has been found to be associated with the lack of IRF1 and STAT1 binding activities as well as of JAK1, JAK2, and STAT1 phosphorylation upon incubation with IFN-γ (13). Although JAK1 and/or JAK2 gene transfer could not restore the IFN-γ-mediated phosphorylation in this RCC cell line, their overexpression increased constitutive LMP2 and TAP1 expression independent of IFN-γ (13). Furthermore, an impaired STAT1 phosphorylation was accompanied by loss of IFN-γ-mediated HLA class I upregulation in melanoma and colorectal carcinoma cell lines (14). The purpose of this study was to determine the mechanisms of IFN-γ unresponsiveness of melanoma cells regarding the HLA class I upregulation as well as the role of the IFN-γ signal cascade for HLA class I APM component expression. Our results show loss of JAK2 expression in 1 of 8 melanoma cell lines, which associated with a lack of IFN-γ inducibility of HLA class I surface antigens and with a low constitutive HLA class I APM component expression. These defects could be corrected by JAK2 transfection; vice versa, JAK2-specific short hairpin RNA and the pharmacologic inhibitor AG490 inversely impairs constitutive APM component expression in JAK2-positive cells, which is associated with reduced HLA class I surface expression.

Material and Methods

Tissue culture

Eight human melanoma cell lines, which have already been described elsewhere or were obtained from the European tumor cell line database (ESTDAB project; see www.ebi.ac.uk/ipd/estdab) were grown in RPMI 1640 medium supplemented with 10% heat-inactivated fetal calf serum (FCS; PAA Laboratories), 2% glutamine (Lonza Cologne AG), and 1% penicillin, and streptomycin (PAA Laboratories) in a humified atmosphere with 5% CO₂.

Antibodies used

The low-molecular-weight polypeptide LMP2, LMP7, and LMP10-specific mouse monoclonal antibody (mAb) SY-1, HB-2, and TO-7, respectively (15), the TAP1-specific mAb NOB-1 (16), the TAP2-specific mAb NOB-2 (16), the tapasin-specific mAb TO-3 (17), and the HLA class I HC-specific mAb HC-10 (18) were developed and characterized as described. All of these are IgG1 mAbs with the exception of mAb HC-10, which is an IgG2a mAb. In addition, following antibodies were purchased, which were directed against the IFN-γR1 (clone 92101; R&D Systems), IFN-γR2 (clone MMHGR-2; Abcam), HLA-ABC (clone B9.12.1; Beckman Coulter), and HLA class II antigens (clone Tū39; Becton Dickinson, BD). The antibodies directed against the unphosphorylated and phosphorylated IFN signal transduction pathway components JAK1 (clone 6G4), pJAK1 (Tyr1022/1023), JAK2 (clone 24B11), STAT1 (clone 42H3), and pSTAT1 (clone 58D6) were all obtained from Cell Signaling (New England Biolabs GmbH). The fluorescein isothiocyanate (FITC)-conjugated IgG2a antibody (Beckman Coulter) served as a control in flow cytometry. The anti-GAPDH (clone 14C10; Cell Signaling) and the anti-β-actin (Abcam) mAbs served as loading controls, whereas the horseradish peroxidase (HRP)-conjugated anti-rabbit and

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anti-mouse IgGs were used as detection antibody in Western blot analysis.

Cytokines and pharmacologic agents
Recombinant human IFN-γ, IFN-α, and TNF-α were purchased from Pan Biotech.

Flow cytometry
The expression of IFN-γR and HLA class I and class II surface antigens was assessed by direct immunofluorescence. For determination of surface expression, 1 x 10⁵ cells were trypsinized, washed with PBS containing 1% FCS, and consecutively incubated with FITC-conjugated respective antibodies for 30 minutes at 4°C in the dark. After 2 washes with PBS, flow cytometry was carried out using the BD FACScalibur flow cytometer (Becton Dickinson). The results were expressed as mean specific fluorescence intensity (MFI) ± SD of the values obtained in 3 independent experiments. Staining with an FITC-conjugated respective IgG antibodies served as the negative control.

Growth properties
A total of 3 x 10⁴ cells were plated in triplicate in the cavities of a 96-well plate overnight before supplementation of different concentrations of IFN-γ (200–1,600 units). After 48 hours, cell viability was measured by standard XTT [2,3-bis(2-methoxy-4-nitro-5-sulfophenyl)-3H-tetrazolium-5-carboxanilide inner salt] method. Results are expressed as relative growth in comparison with untreated cells of 3 independent experiments.

Semi-quantitative RT-PCR and quantitative RT-PCR analysis
For quantitative RT-PCR (qRT-PCR), total cellular RNA was extracted using the RNeasy Mini Kit (Macherey and Nagel), followed by digestion with DNase I (Invitrogen). cDNA was synthesized from 2 μg of total RNA employing the RevertAid H Minus First Strand cDNA Synthesis Kit (Fermentas), according to the manufacturer’s instructions. Comparative quantification of gene expression was carried out by qRT-PCR on a Rotor Gene 6000 system (Corbett Research), using the quantitative SYBR green kit (Invitrogen GmbH) and the target-specific primers listed in Table 1. Amplifications were carried out by an initial hold at 50°C for 2 minutes, followed by denaturation at 95°C for 2 minutes. After 40 cycles with a denaturation step at 95°C for 15 seconds and annealing at 60°C for 30 seconds, the melting steps occurred, starting at 60°C and increasing to 99°C by 1°C steps. For STAT1, the melting step started at 55°C, rising to 99°C. The melting curve analysis was provided at the end of each run to control PCR specificity. Results of the qRT-PCR data were expressed as relative mRNA expression quantified with the Rotor Gene analysis software and normalized to the averaged glyceraldehyde 3-phosphate dehydrogenase (GAPDH) and peptide prolyl isomerase A (PPIA) transcription levels.

For semiquantitative RT-PCR, cDNA was synthesized from 500 ng of total RNA employing the RevertAid™ H Minus First Strand cDNA Synthesis Kit (Fermentas) according to the manufacturers’ instructions. Amplification was carried out in a final volume of 25 μL supplemented with 1.25 units of Taq polymerase, MgCl₂ (Invitrogen), and 0.2 mmol/L dNTP mix (PeqLab). Reaction conditions for the initial denaturation step were 95°C for 2 minutes, followed by denaturation at 95°C for 30 seconds. The annealing occurred at 60°C for 30 seconds, and the extension was done at 72°C for 30 seconds. After 35 cycles, a final extension was done at 72°C for 5 minutes.

Western blot analyses
Proteins (50 μg per lane) extracted from melanoma cells were separated in a 8% to 15% SDS-PAGE gel, depending on the protein size, and transferred to nitrocellulose membranes (Schleicher & Schuell), which were probed with primary mAbs directed against the major HLA class I APM components and molecules of the IFN-γ signal transduction pathway. Following an overnight incubation at 4°C, membranes were incubated for 4 hours with the appropriate HRP-conjugated secondary antibodies as recently described (6). Proteins were detected using a Lumilight detection kit (Roche Diagnostics). Staining of the blots with the anti-β-actin or anti-GAPDH mAbs served as loading controls.

Array comparative genomic hybridization
Genomic DNA was extracted from cultured Colo 857 melanoma cells (test sample) or normal donor peripheral blood mononuclear cells (PBMC; reference sample), using the Qiagen Mini Kit. DNA labeling was conducted using a BioPrime array comparative genomic hybridization (aCGH) genomic labeling kit (Invitrogen). Test and reference DNAs (4 μg of each) were labeled with Cy5 and Cy3, respectively, and cohybridized to the cDNA clone microarray printed at the Infectious Disease and Immunogenetics Section, Department of Transfusion Medicine, Clinical Center, NIH, at 65°C overnight with a configuration of 32 x 24 x 23 spots and contained 17,500 cDNAs (19). Microarray slides were scanned at 10-μm resolution on a GenePix 4000 scanner (Axon Instruments) to obtain maximal signal intensities with less than 0.1% probe saturation.

Genomic PCR
Genomic DNA from melanoma cells was extracted using the Qiagen Mini Kit and genomic PCR was carried out as recently described (20). The forward and reverse primers used for JAK2 amplification were 5’-cat tcc ctt ggg aaa tct ga-3’ and 3’-tgc atg tga aaa cac aca cg-3’, respectively.

Transfection
The JAK2 cDNA was amplified using JAK2-specific primers: forward: 5’-aaa atc gat atg gga atg gcc tgc ct ac-3’ and reverse: 3’- ttg gcc ggc gtc gat cca gcc atg tta ccc c-3’. The 3,339-bp amplification product was directly cloned into...
Duplicate data points were merged by taking the average done using Web tool prep (http://prep.bioinfo.cnio.es/). Tools developed by the Biometric Research Branch, National Cancer Institute (22). Further data preprocessing was performed using BRBArrayTools developed by the Biometric Research Branch, National Cancer Institute (22). Further data preprocessing was done using Web tool prep (http://prep.bioinfo.cnio.es/). Duplicate data points were merged by taking the average done using Web tool prep (http://prep.bioinfo.cnio.es/). Duplicate data points were merged by taking the average done using Web tool prep (http://prep.bioinfo.cnio.es/). Duplicate data points were merged by taking the average done using Web tool prep (http://prep.bioinfo.cnio.es/). Duplicate data points were merged by taking the average done using Web tool prep (http://prep.bioinfo.cnio.es/). Duplicate data points were merged by taking the average done using Web tool prep (http://prep.bioinfo.cnio.es/). Duplicate data points were merged by taking the average done using Web tool prep (http://prep.bioinfo.cnio.es/). Duplicate data points were merged by taking the average done using Web tool prep (http://prep.bioinfo.cnio.es/). The fluorescence intensity and ratio data for Cy5 and Cy3 were transformed into a log base 2 scale and normalized against the median over the entire array using BRBArrayTools developed by the Biometric Research Branch, National Cancer Institute (22). Transfectants were selected in 800 μg/mL G418, and neoR clones were cultivated and further analyzed.

### Table 1. Primers used for RT-PCR analysis

<table>
<thead>
<tr>
<th>Application</th>
<th>Gene name</th>
<th>Sequence</th>
<th>T&lt;sub&gt;m&lt;/sub&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>IFN-γ signaling</td>
<td>JAK1</td>
<td>fw: 5'-gca cca tca ccc tgt atg ac-3'</td>
<td>60°C</td>
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<tr>
<td>rev: 3'-tcc agt gac ctg gca tca ag-5'</td>
<td>JAK2</td>
<td>fw: 5'-tgt gga gat tgt ccc gta tg-3'</td>
<td>60°C</td>
</tr>
<tr>
<td>rev: 3'-att acg ccc acc agc act gt-5'</td>
<td>STAT1</td>
<td>fw: 5'-atc ctc gag agc tgt cta-3'</td>
<td>55°C</td>
</tr>
<tr>
<td>rev: 3'-gcc agq tac tgt ctg att-5'</td>
<td>LMP2</td>
<td>fw: 5'-tgt gca ctc tct gtt tca gc-3'</td>
<td>60°C</td>
</tr>
<tr>
<td>rev: 3'-tgc tgc atc cac ata acc at-5'</td>
<td>LMP10</td>
<td>fw: 5'-ggg ctt ctc ctt cga gaa ct-3'</td>
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<tr>
<td>rev: 3'-tgg ccc cac agc agt at-5'</td>
<td>TAP1</td>
<td>fw: 5'-gga atc ctc tgt ggc aat gtc ca-3'</td>
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</tr>
<tr>
<td>rev: 3'-ttc atc cag cag cac ctg tc-5'</td>
<td>TAP2</td>
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<td>60°C</td>
</tr>
<tr>
<td>rev: 3'-acc tgt cct tgc agg tgt gat-5'</td>
<td>Tapasin</td>
<td>fw: 5'-tgg gta agg gac atc tgc tc-3'</td>
<td>60°C</td>
</tr>
<tr>
<td>rev: 3'-ggt ggc ctc atg gtc aga ga-5'</td>
<td>HLA-ABC</td>
<td>fw: 5'-gcc tac gac ggc aag gat tac-3'</td>
<td>60°C</td>
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<tr>
<td>Locus specificity</td>
<td>RFX3</td>
<td>fw: 5'-aaa ctt gac gca cca gtc aat gc-3'</td>
<td>55°C</td>
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<tr>
<td>rev: 3'-tgt tgt ctc tgt tgt ct-5'</td>
<td>RCL1</td>
<td>fw: 5'-tct tgt tct tgt gtc gca at-3'</td>
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<tr>
<td>rev: 3'-atg ggc ttc aag acc ttc ct-5'</td>
<td>GLDC</td>
<td>fw: 5'-tcg atg cag ttc acc tca ag-3'</td>
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</tr>
<tr>
<td>rev: 3'-cag att ccc acc tga gca tt-5'</td>
<td>BNC</td>
<td>fw: 5'-gtc aag cat gcc tgt gaa g3'</td>
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</tr>
<tr>
<td>rev: 3'-gcc cct ctt tgt tgt cc-5'</td>
<td>GAPDH</td>
<td>fw: 5'-ccc gca cca cca act gct ta-3'</td>
<td>60°C</td>
</tr>
<tr>
<td>Control</td>
<td>PPIA</td>
<td>fw: 5'-cca tgt atg ggg aga aat tgt a-3'</td>
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<tr>
<td>rev: 3'-cag tca gca atg atg atc ttc t-5'</td>
<td>p-Actin</td>
<td>fw: 5'-tgg tgt ggc atc cac gaa act-3'</td>
<td>60°C</td>
</tr>
</tbody>
</table>

Abbreviations: fw, forward; rev, reverse.

The multiple cloning site of the pCMV-IREs expression vector as previously described (21). JAK2-negative cells were stably transfected with the JAK2 expression vector or the control vector (mock), using Lipofectamine (Invitrogen) according to the manufacturer’s instructions. Transfectants were selected in 800 μg/mL G418, and neoR clones were cultivated and further analyzed.

**Data analysis of comparative genomic hybridization (aCGH)**

The fluorescence intensity and ratio data for Cy5 and Cy3 were transformed into a log base 2 scale and normalized against the median over the entire array using BRBArrayTools developed by the Biometric Research Branch, National Cancer Institute (22). Further data preprocessing was done using Web tool prep (http://prep.bioinfo.cnio.es/). Duplicate data points were merged by taking the average over the UniGene cluster IDs. Missing data were imputed using K-nearest neighbor imputation method with K = 15. Gene location was extracted according to UniGene cluster IDs from Ensemble and University of California at Santa Cruz by using ID converter (http://idconverter.bioinfo.cnr.io/). The preprocessed aCGH data were then segmented using the circular binary segmentation method implemented in ADaCGH (http://adacgh.bioinfo.cnio.es/) to detect regions with abnormal DNA copy number. The variables were set as defaults. The significance level for the test to accept change points was set to be 0.01 under 10,000 permutations. The cutoff for gain/loss calls was F0.12 on the log base 2 scale (22). In PBMCs from normal volunteers, this threshold was higher than the 99th percentile of data obtained from autosomes, excluding chromosome X ratios that fell under the 15th percentile. This threshold was then applied to pick up the gain/loss regions...
in the Colo 857 cell line. To ensure the reproducibility of the array data, the array experiments were repeated twice by using DNA isolated from 2 different passages.

Results

**Impaired constitutive HLA class I APM component expression in IFN-γ–resistant melanoma cells**

Flow cytometric analysis of 8 untreated or IFN-γ–treated melanoma cell lines using the HLA class I antigen-specific mAb B9.12.1 (Fig. 1A and C, Supplementary Fig. S1) or the HLA class II-antigen-specific mAb Tu39 (Fig. 1C, Supplementary Fig. S1) showed a marked variability in the IFN-γ–mediated modulation of both HLA antigen classes. The different melanoma cell lines heterogeneously responded in a dose- and time-dependent manner to IFN-γ treatment, ranging from lack of to low to strong IFN-γ–mediated upregulation of HLA class I (Supplementary Fig. S1A) and class II surface antigens (Supplementary Fig. S1B). The representative results shown in Supplementary Figure S1 show that 4 of 8 melanoma cell lines tested exhibited a 2- to 3-fold upregulation of both HLA class I and class II surface antigens, whereas the remaining 4 failed to upregulate HLA class II antigens. The melanoma cell line Colo 857 was completely resistant to IFN-γ treatment, lacking IFN-γ–mediated upregulation of both HLA class I and class II surface antigens (Fig. 1A and B) as well as responsiveness to the antiproliferative effect of IFN-γ (Fig. 1C).
The resistance of Colo 857 cells was selective for IFN-γ because HLA class I surface expression was induced in these cells in a dose- and time-dependent manner by IFN-α as well as by TNF-α, although the degree of upregulation varied between both cytokines (Supplementary Fig. S2). Because the IFN-γ receptor was expressed in the IFN-γ-resistant Colo 857 cells to levels comparable with the IFN-γ-sensitive control cell line Colo 794 (Fig. 1D), the IFN-γ resistance seemed to be due to defects in the IFN-γ signal transduction pathway rather than at the receptor level.

To investigate whether the loss of IFN-γ inducibility of HLA class I surface antigens was associated with altered expression levels of HLA class I APM components, constitutive and IFN-γ inducible LMP10, TAP2, tapasin, HLA class I HC (Fig. 2), LMP2, TAP1, and β2-m (data not shown) mRNA and protein expression levels were monitored by qRT-PCR and Western blot analysis. With the exception of β2-m, the constitutive expression pattern of these molecules was lower and not inducible in IFN-γ-resistant Colo 857 cells than that in IFN-γ-sensitive Colo 794 melanoma cells (Fig. 2). In contrast, IFN-α treatment increased APM component transcription levels and protein expression in both Colo 857 and Colo 794 cells (data not shown).

Lack of IFN-γ sensitivity due to defects in the IFN-γ signal cascade

To determine whether the resistance of Colo 857 cells to IFN-γ is due to an impaired IFN-γ signal transduction, constitutive and IFN-γ-inducible transcription of the major signal transduction molecules including JAK1, JAK2, and STAT1 were investigated. In contrast to the IFN-γ-sensitive cell line Colo 794, RT-PCR revealed a lack of constitutive and IFN-γ-inducible JAK2 mRNA expression in Colo 857 cells, whereas the mRNA of JAK1 and STAT1 was expressed in these cells (data not shown). With the exception of JAK1, the signal transduction components were not upregulated by IFN-γ (data not shown). These
data were further confirmed by Western blot analysis, which showed JAK2 protein expression in Colo 794 and all other melanoma cells analyzed (data not shown) but not in Colo 857 cells despite their constitutive expression of JAK1 and STAT1 proteins (Fig. 3). The functionality of the IFN-γ signaling components was determined using antibodies specifically directed against the selected phosphorylated counterparts JAK1 and STAT1. In Colo 794 cells, an increased phosphorylation of JAK1 and STAT1 was observed after IFN-γ treatment. In contrast, phosphorylation of JAK1 and STAT1 was not detected in Colo 857 cells (Fig. 3). This defect is selective for IFN-γ, as IFN-α did induce STAT1 phosphorylation (data not shown). Thus, the impaired phosphorylation of signal cascade members by IFN-γ treatment reflects the loss of JAK2 expression.

Molecular mechanisms underlying deficient JAK2 expression

To define the molecular mechanisms involved in the lack of JAK2 mRNA and protein expression in Colo 857 cells, JAK2-specific genomic PCR was carried out. Despite the presence of JAK2 amplification products in the Colo 794 control cells, no JAK2-specific genomic PCR product was obtained in Colo 857, suggesting a structural abnormality of JAK2 in these cells (Fig. 4A). In contrast, RT-PCR analyses of 2 genes flanking upstream (RFX3 and RCL1) or downstream (GLDC and BNC2) the JAK2 locus showed amplification products in both the JAK2 + Colo 794 and the JAK2 - Colo 857 cells (Fig. 4B). To confirm these data, CGH of the JAK2 - Colo 857 cell line was done (Fig. 4C) using PBMCs as a control. In addition to multilocus gene amplifications and deletions across the whole genome, a deletion on chromosome 9 from positions 24,466 to 22,022,985 containing the JAK2 gene was found in Colo 857 cells (Fig. 4C). These results indicate that the lack of JAK2 expression in this melanoma cell line is caused by a genomic deletion.

Restoration by JAK2 gene transfer of constitutive and IFN-γ–inducible HLA class I APM component expression by melanoma cells Colo 857

To determine whether the IFN-γ resistance and low levels of HLA class I APM molecules could be restored by JAK2 gene transfer, Colo 857 cells were transfected with a JAK2 expression vector and a control vector (mock) carrying only the neo gene, respectively. The JAK2 transfectants, but not the mock-transfected control cells, did express high levels of JAK2 (Fig. 5A), which was associated with an increased phosphorylation of STAT1 after IFN-γ treatment (Fig. 5A). JAK2 overexpression in JAK2 - cells was accompanied by an enhanced constitutive expression of APM components as representatively shown for TAP1, TAP2, tapasin, and HLA class I HC protein (Fig. 5B), which was accompanied by an increased HLA class I surface antigen expression (Fig. 5C). As expected, the JAK2-transfected cells acquired the susceptibility to modulation by IFN-γ treatment, as indicated by the upregulation of HLA class I APM component expression in these cells (Fig. 5B).

Discussion

The physiologic relevance of the IFN-γ–dependent JAK/STAT pathway was characterized by the functional analysis of JAK/STAT knockout mice and has been linked to antitumor responses (23). IFN-γ can directly act on tumor cells by exerting antiproliferative, proapoptotic, and antiangiogenic effects (24). STATs and JAKs are thought to play a role in promoting these IFN-γ effects on tumor cells, and defects of the JAK/STAT signal transduction intermediates have been associated with an IFN-γ–resistant phenotype in lung carcinoma and melanoma cells (3, 25, 26). This might provide tumor cells with a selective growth advantage. Indeed, more than 30% of human tumors exhibited unresponsiveness or reduced sensitivity to IFN-γ associated with
tumor progression. The variable IFN-γ responsiveness of melanoma cells could be (i) associated with a lower capacity of IFN-γ to induce JAK/STAT (12, 27) or (ii) mediated by downstream components or an additional signaling pathway (28) or (iii) due to lack of STAT1 phosphorylation and epigenetic silencing of the IRF1 transactivation (14). Although abnormalities of HLA class I APM components represent one major mechanism of tumor cells to evade
immune surveillance, there exist only limited information about the role of deficient IFN signal transduction in the regulation of these immunomodulatory molecules. Therefore, the constitutive and IFN-γ-inducible expression pattern of several components of the HLA class I APM and the IFN signal transduction pathway was determined in a number of melanoma cells. With the exception of Colo 857 cells, the other 7 melanoma cell lines analyzed constitutively expressed heterogeneous levels of JAK2 whereas the other IFN signal transduction molecules were constitutively expressed and upregulated by IFN-γ in these tumor cells. Although genetic abnormalities of the IFN-γ signal pathway such as mutations, deletions, and recombinations have been described in tumors of distinct origin (29), no structural alterations in these molecules have been yet reported in melanoma. In this context, it is noteworthy that resistance of melanoma cells to IFN-α is due to multiple defects in the type I IFN signal transduction pathway including lack of Tyk2 (tyrosine kinase 2; ref. 30).

Our results showed that loss of JAK2 expression in the melanoma cell line Colo 857 was caused by a deletion of the JAK2 gene, which is accompanied by a defective IFN-γ signaling and lack of the IFN-γ-mediated HLA class I inducibility. Even more important, impaired JAK2 expression significantly downregulated the constitutive mRNA and protein levels of HLA class I APM components despite a functional APM pathway, thereby providing a selective advantage to tumors. However, this was neither mediated by loss of the IFN-γ R, as Colo 857 cells express this receptor, nor by abnormalities of components of the HLA class I APM. The latter was confirmed by the induction of HLA class I APM molecules, which was accompanied by an upregulation of HLA class I surface expression in these cells on TNF-α and IFN-α treatment, respectively. In addition, JAK2 is required for IFN-γ-induced growth inhibition, as Colo 857 cells lacking JAK2 were not growth inhibited by IFN-γ in contrast to JAK2+ melanoma cells such as Colo 794. This loss of growth-restraining functions might influence tumor progression of JAK2− cells, which will be investigated in future studies.

To confirm the importance of a functional IFN-γ signaling for constitutive HLA class I APM component expression, JAK2 expression was restored in the JAK2− melanoma cells Colo 857 by stable transfection by using a JAK2-specific expression vector. JAK2 gene transfer into JAK2− Colo 857 cells increased the constitutive HLA class I APM component and surface antigen expression. Furthermore, functional JAK2 restored IFN-γ inducibility of HLA class I APM components. Thus, there exists a direct link between abnormalities of HLA class I antigen processing and presentation molecules and impaired JAK2 function. These might also result in reduced CTL sensitivity but increased susceptibility to natural killer cell–mediated lysis. Although a positive correlation between JAK2 and HLA class I antigens has been proven in this study for the first time, a recent publication has shown an improved patient survival when tumors expressed high MHC class I and STAT1 levels in association with a broad T-cell infiltrate (31). Furthermore, loss of STAT1 signaling has been shown to be associated with a higher incidence of tumors in mice (23, 32). These results strengthen our hypothesis of an important role of a functional IFN-γ signal cascade for the immunosurveillance of tumor cells.

Owing to the complexity of the IFN-γ signal transduction pathway, a comprehensive explanation how and at which level other elements of the IFN-γ system besides JAK2 and STAT1 modulate HLA class I APM component expression is still awaiting. Because JAK2 is a key regulator of IFN-γ responses and is induced by other growth factors and DNA damage, tumors acquiring resistance to IFN-γ by dysregulation or structural alterations of JAK2 might evade the immunosurveillance leading to tumor progression; vice versa, an impaired IFN-γ signaling in association with a reduced HLA class I APM component expression pattern suggests that defects in the IFN-γ
cascade might play a crucial role in the malignant transformation process and might be involved in the frequent development of immune escape phenotypes caused by HLA class I APM component abnormalities. This is further supported by cDNA microarray analysis of different tumors showing an altered expression of IFN signaling molecules. In conclusion, the present study identified for the first time that a deletion of JAK2 in melanoma cells or inhibition of JAK2 signaling caused an impaired MHC class I APM component expression, suggesting that abnormalities of the JAK/STAT pathway might play an important role during development of melanoma. In addition, these results underscore the biological significance of JAK2 for IFN-γ treatment-independent immunosurveillance and provide evidence that defects in JAK2 might represent a novel immune escape mechanism and a potential target in combination with T-cell–based therapies. Further experiments are currently been carried out to validate these data in a large series of tumor lesions by using a tissue microarray and associate them with the survival of melanoma patients.

Disclosure of Potential Conflicts of Interest

The authors have no conflicting financial interests.

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Disclosure of Potential Conflicts of Interest

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