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Survival of metastatic melanoma patients after dendritic cell vaccination correlates with expression of leukocyte phosphatidylethanolamine-binding protein 1 / Raf Kinase inhibitory protein

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ABSTRACT

Immunotherapy for metastatic melanoma offers great promise but, to date, only a subset of patients have responded. There is an urgent need to identify ways of allocating patients to the most beneficial therapy, to increase survival and decrease therapy-associated morbidity and costs. Blood-based biomarkers are of particular interest because of their straightforward implementation in routine clinical care. We sought to identify markers for dendritic cell (DC) vaccine-based immunotherapy against metastatic melanoma through gene expression analysis of peripheral blood mononuclear cells. A large-scale microarray analysis of 74 samples from two treatment centers, taken directly after the first round of DC vaccination, was
performed. We found that phosphatidylethanolamine binding protein 1 (PEBP1)/Raf Kinase inhibitory protein (RKIP) expression can be used to identify a significant proportion of patients who performed poorly after DC vaccination. This result was validated by q-PCR analysis on blood samples from a second cohort of 95 patients treated with DC vaccination in four different centers. We conclude that low PEBP1 expression correlates with poor overall survival after DC vaccination. Intriguingly, this was only the case for expression of PEBP1 after, but not prior to, DC vaccination. Moreover, the change in PEBP1 expression upon vaccination correlated well with survival. Further analyses revealed that PEBP1 expression positively correlated with genes involved in T cell responses but inversely correlated with genes associated with myeloid cells and aberrant inflammation including STAT3, NOTCH1, and MAPK1. Concordantly, PEBP1 inversely correlated with the myeloid/lymphoid-ratio and was suppressed in patients suffering from chronic inflammatory disease.

INTRODUCTION

Metastatic melanoma is one of the most devastating types of cancers in terms of potential life-years lost and affects a growing number of patients each year [1, 2]. Currently, there are different types of immunotherapy that may offer long-term benefit for melanoma patients. These therapies are based on: 1) antagonists of molecules that suppress pre-existing anti-tumor immune responses, so-called immune checkpoint inhibitors such as anti-CTLA-4 (ipilimumab) and anti-PD-1/PD-L1 antibodies (nivolumab/pembrolizumab/atezolizumab); 2) delivery of autologous ex vivo expanded tumor infiltrating lymphocytes to boost anti-tumor T cell response; 3) oncolytic viruses injected into metastases to lyse tumor cells and enhance immune responses; and 4) ex vivo or in vivo targeting of dendritic cell (DCs) to initiate and/or boost tumor antigen-specific immune responses (reviewed in [3, 4]). These new immunotherapies, however, are extremely expensive and/or labor intensive due to high production costs and/or the need for patient-personalized preparation. In addition, although these therapies offer clear benefit for a group of patients, there are also many primary and secondary non-responders [5–7]. This is illustrated by the fact that individual melanoma patients can respond very well to immunotherapy, but those that do represent a limited proportion of the total number who receive this treatment [8–13]. Markers to determine which patients do, or do not, benefit are therefore urgently needed to facilitate treatment decisions. In particular, mechanism of action-based markers, derived from our understanding of why specific treatments are beneficial or not in certain patient populations, have a high potential [7].

So far, a number of prognostic factors for survival in melanoma have been defined, together with several markers associated with or predicting responses to various forms of therapy [14–17]. The success of many immunotherapies is thought to be associated with the composition and organization of the tumor microenvironment, and therefore biomarkers are often sought in this respect [18]. Indeed several studies have identified tumor properties that have correlated the with outcome of immunotherapies, including the expression of cytokines (interferon-γ), chemokines (CCL5, CXCL9, CXCL10), immunoregulatory molecules such as indolamine-2,3-dioxygenase [8, 19] and, more recently, tumor expression of PD-L1 that was reported to be predictive for responses to checkpoint inhibitors in melanoma [20].

Many of the potential biomarkers for immunotherapy against melanoma, reported so far, have been tissue based. This severely limits their clinical applicability because for many patients the primary tumor is not or no longer available, and metastatic tumor tissue material can also be hard to obtain. Furthermore, intra-individual heterogeneity of tissue biomarker expression (i.e. between different metastases and/or with respect to the primary tumor) may limit reliability [21]. These factors can severely limit the predictive effectiveness of a biomarker and, more importantly, may lead to the incorrect exclusion of patients from a potentially effective therapy. Furthermore, tissue biomarkers can only be assessed at one single time point. In contrast, blood-based markers can potentially overcome all these issues. Diagnostic liquid biopsies have already been used in breast cancer, colorectal cancer and in non-small cell lung cancer [22–24]. Moreover, blood samples are being utilized for cell-based cancer-detection in plasma and for tumor mutational analyses [24–26]. A major advantage of obtaining blood samples as a source of biomarkers for therapy prediction is that it is a minimally-invasive intervention which allows the tracking of disease dynamics during the course of treatment.

For dendritic cell DC vaccination specifically, efforts have been made to identify markers associated with patient responses to therapy [27, 28]. In our laboratory, we have developed a delayed type hypersensitivity response (DTH) test to assess the induction of tumor-specific T cells in response to DC vaccination into the skin [29, 30]. This showed that the induction of antigen-specific T cells by DC vaccination was associated with improved survival. However the absence of detection of tumor-reactive T cells did not necessarily indicate a poor response, and therefore additional methods to predict whether a patient
will benefit from DC therapy are required. Here we have used microarray analysis of peripheral blood mononuclear cells (PBMCs) from treated patients to identify biomarkers that can be used to monitor and further improve our understanding of the response to DC vaccination. Using this strategy we have identified and validated phosphoethanolamine binding protein 1 (PEBP1), also known as Raf-1 kinase inhibitor protein (RKIP), as a biomarker that can be used to predict patient survival after DC vaccination. PEBP1 has previously been described to modulate several major cancer and inflammatory signaling pathways [31–34]. Here, further in-depth analysis of the genes that are co-expressed with PEBP1 suggests that its increased expression after vaccination is indicative of the beneficial skewing of the adaptive immune system towards an effective anti-tumor response.

RESULTS

Selection of genes with prognostic potential from microarray data of PBMCs from treated patients

To identify biomarkers in the blood of treated patients that may correlate with survival after DC vaccination, we analyzed PBMCs from a total of 74 patients by microarray (MA). These patients had been treated with DC vaccination in two different centers (Discovery cohort; Nijmegen and Erlangen). Only PBMC samples taken after vaccination were at our disposal in sufficient numbers to allow MA-based comparison of good and poor responders. In search of a reasonably-sized set of potential biomarkers that allowed validation by qPCR, we performed several different statistical analyses, including survival analysis, statistical analysis of microarray (SAM) as well as multivariate analysis (PLS-DA; Supplementary Figure 1). For survival analysis, expression data were related to survival time after the start of DC vaccination in a continuous manner. For SAM and PLS-DA, patients were grouped into three survival subgroups [short (<1 year); medium (1-2 years), and long (>2 years); Table S1], to allow us initially to search for gene expression differences between the extremes of this survival spectrum. For SAM, we extracted probes with expression values satisfying two criteria: (i) significant differences between patients from the short and long survival subgroups; and (ii) significant differences between PBMCs and vaccine DCs of the same donor (to increase the chance of finding biomarkers related to a specific cell type and/or immune response). For all three methods, a final filter taking at least a 2-fold change in expression levels between the short and long survival subgroups was applied (Supplementary Figure 1). These three approaches resulted in relatively large lists ranging from 62 to 307 candidates that all may have potential prognostic value. To further reduce the list to a size compatible with qPCR validation, only the candidates that emerged from all three approaches were retained. This final list consisted of 19 genes (Table 1).

Validation of microarray analysis results by qPCR

Next, we set out to verify MA results using qPCR. For MA analysis we had chosen to use as many datasets as possible in order to find the most robust differentially-expressed genes across treatment centers, and to add power to the statistical analysis. This strategy, however, posed the risk of a center effect since the samples from the two treatment centers were not fully comparable in terms of patient composition and timing of sample collection (methods and supplementary tables S1-S3). In addition, while stage IV patients comprised the majority, stage III melanoma patients had also been included (Supplementary table S1 and S2). Thus, to exclude any confounding effects of these dissimilarities, only Nijmegen-center stage IV patients were used for the initial qPCR validation of the MA result and for the subsequent selection of qPCR validated genes. For 28 patients within this group, we first directly compared the MA data to the qPCR data obtained, to assess which genes behaved most consistently using these very different techniques. Out of 19 genes derived by MA, 11 showed a significant and positive correlation with the qPCR measurement (Supplementary information Figure 2 & Table 1).

We then assessed whether the expression of any of the above qPCR-validated genes also correlated with survival. After correction for multiple testing, four genes out of the 11 passed this criterion (Supplementary information Figure 3 & Table 1). Only one of these genes, PEBP1 (RKIP) correlated positively with patient survival; the remaining three, FCGR1B (CD64), ANXA2 (Annexin-2) and FCGR1B (Fcγ-receptor 1B, CD64), correlated negatively (Table 1). Hence these four genes could potentially predict clinical outcomes and/or provide mechanistic insights into the mechanisms of patient responses to DC therapy.

Validation of differential expression of selected genes between patient groups by qPCR

We wished to further assess and validate the association of the above 4 genes with patient survival after DC vaccination. Therefore we tested their expression on a second novel set of samples derived from Nijmegen patients that had not been used for MA gene selection, and on a completely independent set of samples derived from Copenhagen, a third treatment center exploiting DC vaccination. Interestingly, for PEBP1 the expected positive correlation with survival time, as well as a significantly lower expression in PBMCs from short surviving patients,
was observed in validation samples from both treatment centers (Figure 1). Also, additional samples not used for MA from the Erlangen treatment center showed a similar trend for PEBP1 despite the fact that the timing of sample collection was very different from the other two treatment centers (Supplementary Figure 4, methods). Unfortunately, insufficient samples from short survivors were available from Erlangen to draw any firm conclusions. For FCGR1B, a trend for negative association with survival was observed, but only in the Nijmegen validation cohort (Figure 1). However, the association of ANXA2 and IMPA2 could not be confirmed in any of these samples: they were neither differentially expressed between long and short survivors in the additional Nijmegen samples, nor in the completely independent Copenhagen samples (Figure 1A and data not shown).

### Table 1: Validation of MA expression data by qPCR

<table>
<thead>
<tr>
<th>Gene (Symbol)</th>
<th>Pearson correlation MA / qPCR</th>
<th>p-value</th>
<th>Spearman correlation to survival</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>FCGR1B</strong></td>
<td>0.68</td>
<td>&lt; 0.0001</td>
<td>-0.52</td>
<td>0.0023</td>
</tr>
<tr>
<td><strong>ANXA2</strong></td>
<td>0.47</td>
<td>0.0059</td>
<td>-0.46</td>
<td>0.0065</td>
</tr>
<tr>
<td><strong>PEBP1</strong></td>
<td>0.46</td>
<td>0.0064</td>
<td>0.46</td>
<td>0.0074</td>
</tr>
<tr>
<td><strong>IMPA2</strong></td>
<td>0.45</td>
<td>0.0086</td>
<td>-0.34</td>
<td>0.0407</td>
</tr>
<tr>
<td><strong>MNDA</strong></td>
<td>0.58</td>
<td>0.0006</td>
<td>-0.31</td>
<td>0.055</td>
</tr>
<tr>
<td><strong>SLC4A7</strong></td>
<td>0.49</td>
<td>0.004</td>
<td>0.29</td>
<td>0.0679</td>
</tr>
<tr>
<td><strong>BST1</strong></td>
<td>0.45</td>
<td>0.0078</td>
<td>-0.26</td>
<td>0.0898</td>
</tr>
<tr>
<td><strong>MGST1</strong></td>
<td>0.48</td>
<td>0.0048</td>
<td>-0.24</td>
<td>0.1074</td>
</tr>
<tr>
<td><strong>ZNF467</strong></td>
<td>0.38</td>
<td>0.0244</td>
<td>0.09</td>
<td>0.3288</td>
</tr>
<tr>
<td><strong>OAS2</strong></td>
<td>0.42</td>
<td>0.0136</td>
<td>-0.02</td>
<td>0.4637</td>
</tr>
<tr>
<td><strong>UBE2L6</strong></td>
<td>0.33</td>
<td>0.0428</td>
<td>0.02</td>
<td>0.4681</td>
</tr>
<tr>
<td><strong>ATP1B3</strong></td>
<td>0.20</td>
<td>0.1505</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td><strong>FH</strong></td>
<td>0.16</td>
<td>0.2113</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td><strong>LILRA5 (primer 1)</strong></td>
<td>0.31</td>
<td>0.0527</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td><strong>LILRA5 (primer 2)</strong></td>
<td>0.28</td>
<td>0.0779</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td><strong>TPM3 (probe 1)</strong></td>
<td>-0.16</td>
<td>0.2147</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td><strong>TPM3 (probe 2)</strong></td>
<td>-0.10</td>
<td>-0.0967</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td><strong>SRSF6</strong></td>
<td>0.01</td>
<td>0.4705</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td><strong>MAT2B</strong></td>
<td>-0.04</td>
<td>0.4255</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td><strong>PSMB9</strong></td>
<td>-0.11</td>
<td>0.2933</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td><strong>TLR6</strong></td>
<td>0.14</td>
<td>0.2389</td>
<td>NA</td>
<td>NA</td>
</tr>
</tbody>
</table>

For the total 28 samples from stage IV patients of the Nijmegen discovery cohort we calculated the correlation between the expression levels as measured by microarray analysis (MA) against those obtained by qPCR. For validated genes we also calculated the correlation between qPCR expression and survival time. Only FCGR1B, ANXA2, PEBP1 and IMPA2 were validated (in bold). a Two different primers were tested. b Two probes for this gene were put forward by our statistical pipeline. See also Supplementary Figures 2 and 3 for correlation plots. NA: Not analyzed.

### Assessment of the clinical value of PEBP1 as a biomarker

Having established that the patients performing poor after DC vaccination expressed lower levels of PEBP1 in PBMCs acquired after vaccination, irrespective of the treatment center, we set out to quantify the efficacy of PEBP1 to discriminate short- and long-surviving patient populations. To this end, we first performed a normalization step based on the median value of PEBP1 measured on all samples from each center and subsequently merged all Nijmegen and Copenhagen data (Figure 2; Table S1 for patient numbers). We then correlated the normalized expression data from all patients with survival time. On this merged qPCR dataset, containing all stage IV patient samples from Nijmegen and
Copenhagen, we again observed a significant correlation of PEBP1 expression with survival, confirming the potential of PEBP1 as a biomarker (Figure 2A). By visualizing the total patient population in one plot, it became apparent that, while most well-responding patients displayed a high expression of PEBP1, poorly responding patients could be subdivided into two groups with either low or high PEBP1 expression. This indicates that while low PEBP1 expression for the vast majority of patients marks a poor clinical outcome after DC vaccination, high PEBP1 expression does not necessarily indicate a good prognosis. Nevertheless, a low expression of PEBP1 after the first

![Figure 1: Validation on independent samples.](image)

Samples from Nijmegen (Nij) that had not been part of the microarray (MA) study (A) and samples from a third completely-independent treatment center, Copenhagen (Cop; B), were tested to validate the MA-predicted association with patient survival. Upper panels: expression (qPCR) versus survival time. Spearman r and p-value (one-tailed) are given. Lower panels: expression (qPCR) of indicated genes in short (<1 year) versus long (>1 year) surviving patients in the two treatment centers. p-value by t-test (one-tailed).
A set of vaccinations could identify potential non-responders early-on, and thus enable early change to alternative or additional treatment. To find the optimal cut-off level of \textit{PEBP1} expression and survival length to identify such patients, we generated receiver-operator curves (ROC). The ROCs were based on the predetermined 12-month cut-off (i.e., the short survival group). This ROC performed well [area under the curve (AUC) 0.67+/-0.053]. The correlation plot, however, indicated that a survival of 14 months rather than 12 months best delineated the patients with low \textit{PEBP1} expression and low survival. Indeed, patient selection was further improved when the maximal survival time for the short survival group was extended to 14 months (Figure 2A). This yielded the largest differences between survival groups and a ROC curve with an AUC of 0.69+/-0.053 (Figure 2B & 2C).

\textbf{Figure 2: Expression of \textit{PEBP1} is of value for treatment decision.} (A) Spearman correlation of center-normalized expression values for \textit{PEBP1} with survival time after DC vaccination (one-tailed p-value). Grey area: population of patients with low expression of \textit{PEBP1} (<-0.33) that have predominantly a limited survival. Specifically indicated are the original 12-month (dotted line) and the adjusted 14-month (broken line in bold) survival windows (B) Center-normalized expression of \textit{PEBP1} in all evaluated patient samples from Nijmegen and Erlangen (discovery and validation together; one-sides p-values by t-test). (C) Receiver Operator Curve (ROC) of all Nijmegen and Copenhagen samples combined to select short survivors based on low expression of \textit{PEBP1}. (D) Kaplan-Meyer curve of patients having a \textit{PEBP1} expression level below or above -0.33 (2 log center normalized C(t)).
Of note, similar or better results were obtained for each treatment center separately, yielding AUC of 0.67+/-0.06 and 0.76+/-0.08 for Nijmegen and Copenhagen respectively (Supplementary Figure 5). Also, the Erlangen dataset showed the same trend (AUC 0.64+/-0.14) but the ROC result was not significant, likely because of the low number of short surviving patients in this cohort (7 patients). Based on the ROC curve of all pooled samples we determined that a PEBP1 expression cut-off of -0.33 (center normalized log(Ct)) allowed us to identify patients who were surviving less than 14 months after vaccination with a sensitivity of 46% and a specificity of 89%. Using this expression cut-off we were thus able to select a patient population with a significantly poorer clinical outcome after vaccination, that is unlikely to respond to further DC therapy in the applied format and which would be in immediate need of other or additional treatments (Figure 2D).

**PEBP1 expression during the course of vaccination**

We then investigated whether PEBP1 expression levels could be informative also before vaccination, and therefore studied the expression of PEBP1 over time. For this we used Copenhagen samples for which PBMCs were taken a few days before vaccination, after 4 vaccinations and after 6 vaccinations. Intriguingly, there were no differences in the expression levels of short and long surviving patients in PBMCs before vaccination, indicating PEBP1 expression correlates with clinical outcome after vaccination but cannot a priori predict whether a patient will respond or not (Figure 3A & 3B). The difference in PEBP1 expression was further increased after two additional vaccinations (Figure 3B & 3C). Notably, we found that the change of PEBP1 in response to vaccination correlated better with survival time than the expression level of PEBP1 itself (Figure 3D). Interestingly, an effect of vaccination was seen both in the short- and long-surviving patients: in short-surviving patients PEBP1 levels decreased during the course of vaccination, whereas it increased in long survivors (Figure 3C). A similar trend was also present in a few longitudinal samples available from a fourth, independent treatment center in Innsbruck (Figure 3E). Together these results suggest that the change in PEBP1 expression (change) is indicative of patient responses to DC vaccination using overall survival as an endpoint, and as such may provide a mechanistic insight into what determines a successful treatment response.

**PEBP1 may indicate skewing of the immune response**

To obtain further insights into how PEBP1 expression may reflect the response to DC vaccination, we interrogated the original MA data to identify genes that were positively or negatively correlated with PEBP1 to understand more about the molecular context in which this protein may act. First, we analyzed genes ranked on their correlation with PEBP1, for the enrichment of previously described blood transcriptional modules (BTMs) using Gene set enrichment analysis (GSEA) [35–37]. Interestingly, BTMs indicated that PEBP1 expression correlated well with genes associated with an adaptive immune response (e.g. T cells and their activation) as well as the regulation of transcription (Table 2; Figure 4). However, the negatively-correlated genes were enriched for those relating to myeloid cells (e.g. monocytes, neutrophils, and immature DCs) and inflammation.

In the search for more mechanistic insights, we subjected the genes that were positively- or negatively-correlated with PEBP1 to protein-protein-interaction and gene ontology (GO) analyses. We found that genes correlating with PEBP1 expression mapped to extremely well-connected, mostly ribosomal, proteins, suggesting high protein synthesis (ability) in these patient samples. Negatively-correlating genes were clustered around the signaling proteins STAT3, NOTCH1, MAPK1 and UBB (Figure 4). Moreover, these genes mapped to GO terms relating to antigen processing and presentation and the response to stress (supplementary Table S5), and in addition to the innate immune response as previously also indicated by GSEA. Together these results indicate that low PEBP1 may represent a state of systemic inflammation and stress accompanied by low protein biosynthesis. This systemic state may hamper an effective response to DC vaccination.

To investigate whether these functional implications of PEBP1 expression may stretch beyond DC vaccination and/or metastatic melanoma, we interrogated a large co-expression database of immune cells (immune-navigator) and found that also in this database PEBP1 expression correlated with adaptive immune cell responses and anti-correlated with monocytes and generic immune activation (Figure 4 and supplementary table S6). Moreover, further analysis of publicly available blood or PBMC expression data comparing healthy subjects to those suffering from chronic inflammatory diseases (e.g Ankylosing spondylitis, Schnitzler syndrome and rheumatoid arthritis) revealed that low PEBP1 expression may be a more frequent event associated with chronic inflammation than previously anticipated (Figure 4).

These data suggest that low PEBP1 expression after DC vaccination may reflect the ineffective development of a productive, adaptive immune response and instead may be associated with myeloid cells that can negatively regulate immune responses in a state of aberrant/chronic inflammation. Recently, several studies have reported the deleterious effect of myeloid-derived suppressor cells (MDSCs) in melanoma and in response to DC vaccination [15, 17, 38, 39]. Indeed, we found that biobanked PBMCs...
Figure 3: Change in PEBP1 expression correlates with survival after vaccination. (A-D) Expression of PEBP1 was assessed in Copenhagen patients prior to or after 4 or 6 rounds of vaccination. (A) Spearman correlation of PEBP1 at the 3 time-points (p-value one-tailed). Respectively n=39, n=38 and n=30 pre-, after 4 and after 6 vaccinations. (B) Expression of PEBP1 at each time-point in short (<14 months) or long survivors (> 14 months). Ns= non-significant * p<0.05 ** p<0.01 by t-test (one-tailed). (C) Course of PEBP1 expression during successive rounds of vaccination in short and long survivors (Mean +/- SEM). (D) Spearman correlation of survival time with the change in PEBP1 expression with respect to the start of vaccination (p-value one-tailed). E Change in PEBP1 expression in relation to survival in patient samples from Innsbruck (n=7 stage IV in black; n=2 stage III in red).
from DC-vaccinated patients demonstrated a trend for the inverse relation between *PEBP1* expression levels and MDSC frequencies in the Nijmegen samples. However, this relationship could not be confirmed for samples of the other centers, possibly due to the non-optimal conditions induced by sample freezing and differences in the MDSC identification strategy (Supplementary Figure 6). In search for an alternative way to further substantiate the notion that high *PEBP1* expression may associate with high lymphocyte/adaptive immune responses but low

Figure 4: *PEBP1* expression reflects metabolic and inflammatory state. (A and B) Examples of BTMs that were found enriched by GSEA in the genes correlating to *PEBP1* in the microarray dataset. Genes positively correlated to *PEBP1* (A) were enriched in BTMs related to T cell (activation), whereas those negatively correlating with *PEBP1* were enriched in BTMs related to monocytes and general inflammation (A). (C and D) Protein-protein Interactions (PPI) present in the STRING database for the top 500 genes positively (C) or negatively correlating to *PEBP1* (D). Shown are genes for which the STRING database reported an interaction with a high confidence score (0.7) and more than one connection. Gene symbols are only displayed for genes with at least 20 (left graph) or 10 connections (right graph). (E) Enrichment of genes from the T cell and monocyte module within a list of genes ranked on correlation with *PEBP1* in the Immuno-Navigator database. (F) Expression of *PEBP1* in blood or PBMC samples of healthy donors versus patients suffering from Ankylosing spondylitis (AS; GDS5231), Schnitzler syndrome (SS; GSE70019) and rheumatoid arthritis (RA; GDS3794) as retrieved using GEO2R.
myeloid/innate immune response we re-analyzed all available flow cytometry data of our patients acquired at the same moment of mRNA sample collection to quantify myeloid and lymphoid cell fractions. In agreement with the GSEA result PEBP1 anti-correlated (or showed this trend) with the myeloid/lymphoid balance in all centers tested (Figure 5). This is further supported by the observation that, a change in PEBP1 expression as a result of vaccination was significantly more often associated with a change of the myeloid/lymphocyte balance in the opposite direction (p<0.01 by Fisher exact test; Figure 5).

Together, these data demonstrate that in patients doing poorly after DC vaccination, low PEBP1 levels are indicative of a distorted myeloid/lymphocyte balance and that this may correlate with skewing of the induced immune response towards chronic inflammation.

**DISCUSSION**

We here describe PEBP1 as a novel gene whose low expression on blood leukocytes is associated with poor survival in metastatic melanoma patients receiving DC therapy. The association of PEBP1 with survival provides important clues towards the mechanisms that determine the success of the DC vaccine-driven immune response. Furthermore, PEBP1 expression on white blood cells may have clinical value in selecting patients who may not benefit from further DC treatment in its current form and who might be better treated with other forms of immune or combination therapy (for example in combination with immune checkpoint blockade or chemotherapy).

We tested and validated the association of PEBP1 with survival on independent samples from different treatment centers. Our data indicate that low PEBP1 levels after DC vaccination, but not before, are associated with a poor survival. Interestingly, we found that in patients who responded poorly, PEBP1 levels decreased even further, whereas patients who did well demonstrated a rise in PEBP1 levels upon vaccination. Based on the molecular context in which we find PEBP1 to be expressed, we postulate that a decrease in PEBP1 may be indicative of a skewing of the (DC-vaccine-triggered) immune response towards chronic inflammation/myeloid immune suppression rather than towards an effective anti-tumor response. Such a scenario would be in line with both the requirement for a robust adaptive response to combat the
tumor, and the reported high prevalence of inflammation and/or tumor-induced MDSCs in patients with a poor prognosis [15-17, 38, 39]. Indeed, in the Nijmegen cohort, PEBP1 levels were negatively correlated with monocytic MDSC numbers. These experiments that aimed to determine the relationship between PEBP1 and MDSC are however by no means conclusive, and were hampered by great variability in MDSCs numbers between centers. The use of frozen material and the non-standardized MDSC identification methods used limited these analyses [40].

Figure 5: Blood PEBP1 expression anti-correlates with the myeloid/lymphoid balance. (A) Spearman rank correlation of PEBP1 expression with the myeloid/lymphocyte (M/L) balance determined by flow cytometry at the time of sample collection for all four treatment centers. (B) Quantification of the direction of change of PEBP1 with respect to the M/L balance in sequential longitudinal samples from Copenhagen (see also figure 3).
Going forward these analyses should be included in future studies. Intriguingly, and in agreement with the reported function and ontogeny of genes correlating and anti-correlating with PEBP1, we found that PEBP1 expression levels in patient samples from all centers was inversely correlated with the myeloid/lymphoid balance in PBMCs at the time of sample collection.

It remains to be determined whether PEBP1 is directly driving the skewing of T cell-mediated immunity towards robust anti-tumor responses, or is merely a bystander gene reflecting the state of the immune system. PEBP1, also known as RKIP, is a mostly-cytoplasmic protein that is widely expressed and has diverse functions affecting several important signaling cascades including NFκB, MAPK/ERK and GSK3β signaling, to regulate cell proliferation, migration, and activation (reviewed by [34, 41]). In cancer, PEBP1 expression seems mostly favorable to survival and, for melanoma, an inverse correlation between PEBP1 expression and metastases has been reported [31, 32]. Although we discovered a similar favorable association between high PEBP1 expression and survival, our finding is on white blood cells, not on tumor cells. Our results are therefore unlikely to be explained by an intrinsic effect of PEBP1 on tumor cells or their metastatic counterparts. Instead it is more likely that PEBP1 acts on the white blood cells themselves. This notion is supported by the inverse correlation of PEBP1 expression with the myeloid/lymphocyte balance. Intriguingly, the genes inversely correlated with PEBP1 included, and were centered around STAT3, NOTCH1, and MAPK1 (ERK2) (by PPI). Possibly the discriminative power of PEBP1 demonstrated here may relate to the signaling pathways associated with these molecules. PEBP1 expression has been reported to dampen the inflammatory response, acting on both ERK/MAPK and NFκB signaling to reduce pro-inflammatory cytokine release [33, 34, 42, 43]. High PEBP1 expression has also recently been shown to attenuate STAT3 signaling, which in turn has been shown to be a key factor for the suppressive effect of MDSCs [44–46]. MDSCs are postulated to dampen the immune response to tumors, including melanoma, and are associated with poor survival [38, 39, 47, 48]. The presence of chronic inflammation in cancer patients has been held responsible for the induction of MDSC via growth factors and cytokines (e.g. G- or GM-CSF, IL-6, TGFβ, PGE2) acting via ERK/ mTOR, STAT, NFκB and SMAD signaling pathways [49, 50]. Considering the inverse correlation between PEBP1, STAT3 and ERK expression identified here, it seems quite possible that PEBP1 acts on these signaling pathways to prevent/dampen excessive systemic inflammation that may hamper an effective adaptive immune response following DC vaccination. This hypothesis is supported by our finding that PEBP1 levels are also diminished in patient suffering from other diseases associated with chronic inflammation.

Besides regulating inflammation in general, the action of PEBP1 may be more sophisticated. In CD8+ T cells PEBP1 expression is associated with enhanced IFN-γ signaling downstream of the TCR [51]. PEBP1 may thus have dual functions in inhibiting deleterious systemic inflammation that drives immunosuppression, whilst at the same time promoting adaptive responses. In line with this view, we find in our gene expression data an inverse correlation of PEBP1 with genes connected to innate and inflammatory responses, while genes associated with T cell responses positively correlate with PEBP1 expression. Furthermore, we observed that PEBP1 is also highly correlated with the expression of ribosomal subunits suggesting PEBP1 expression to be associated with high protein biosynthesis capacity. This may facilitate an effective adaptive immune response which is highly dependent on the production of large amounts of inflammatory mediators (e.g. cytokines, antibodies) and immune cell expansion. Further research on the expression and function of PEBP1 in different immune cells is required to shed further light on how exactly PEBP1 relates to the skewing of immune responses following DC vaccination.

Our study once more indicates that the presence of excessive/chronic inflammation and/or a lack of “fitness” of the immune system may be a severe obstacle to the success of DC vaccination. Suppressive and exhausted immune cells may hamper the effectiveness of treatment modalities that are aimed at inducing or boosting adaptive immune responses. Together, the data presented here and from other studies, suggest that the presence of chronic inflammation, MDSCs and now also low PEBP1 expression may all represent contraindications for further DC vaccination in its current form. More studies are required to obtain detailed mechanistic insight into the interaction of these factors. Recent literature suggests that lifting this state of aberrant inflammation and concordant suppression of immune responses in cancer patients, by e.g. interfering with STAT signaling or by certain regiments of chemotherapy could increase the effectiveness of various types of immunotherapy [49, 52, 53]. As such PEBP1 could represent a facile and effective blood-based biomarker to monitor whether the fitness of the immune system is adequate for successful immunotherapy.

MATERIALS AND METHODS

Patients

Clinical data and biological samples from melanoma patients undergoing DCs vaccination were obtained from the University of Erlangen, Erlangen, Germany (see Table 1; www.clinicaltrials.gov NCT00053391 and [54, 55], from the Radboud University Nijmegen Medical Centre, Nijmegen, The Netherlands (NCT00243594; NCT00243529;
were reverse transcribed to synthesize first- and second-strand complementary DNA (cDNA), purified and in vitro transcribed to synthesize biotin-labeled complementary RNA (cRNA) using the Illumina TotalPrep-96 RNA Amplification Kit (# 4393543 Ambion, Inc., Austin, TX). A total of 750ng of biotin-labeled cRNA was then hybridized to Illumina HumanHT-12 V3.0 expression BeadChips at 55°C for 18 h. The hybridized BeadChip was washed and stained with streptavidin-Cy3 according to the manufacturers protocols using Illumina whole-genome gene expression direct hybridization assay (#11286340 Illumina, San Diego, California, USA). GenomeStudio Data Analysis Software used to visualize and analyse images generated. This software provides data in standard file formats that can be readily processed with most commercial and/or public accessible gene expression analysis software programs to identify significantly differentially expressed genes with pathways and networks involved. Raw data from BeadStudio were exported as text files and processed using the Bioconductor R package Lumi [62]. Quality controls were performed, and samples with incorrect parameters were discarded. The background was adjusted by subtracting an offset estimated based on the quantile of the control probes; then normalization was performed using log2 transformations. All further statistical analyses, unless otherwise specified, was performed on probe expression values. Raw data as well as normalized data for all the samples (74 PBMC samples and 68 matched mature moDCs samples) we used in this work have been deposited in the ArrayExpress repository with the ID E-MTAB-5201.

Microarray data statistical analysis

Survival analysis by SAM

The significance analysis of microarrays (SAM) as implemented in the Bioconductor R package Rsam [63] includes a procedure for estimating survival associated probes given a vector of survival months and the censoring status of the PBMC donor. We used a total of 74 PBMC samples to draw the survival analysis, keeping the 90th percentile FDR = 0 to exclude the presence of false positive survival associated probes. The resulting probes were then associated with a fold change by dividing their mean expression values in long survivors by their mean expression value in short survivors. A fold-change cut-off of 2 was then applied to restrict the number of probes.

Partial Least Square Discriminant Analysis/ Multivariate analysis

Gene expression was also investigated through the multivariate methods implemented in the R package mixOmics [64]. In particular, we used a partial least square analysis (PLS) coupled to a discriminant analysis (DA) to accommodate a two or three-class categorization of the samples. The categories we used were the above...
mentioned ranked survival classes of long, medium and short survivors. Fold changes were calculated as described for the survival analysis by SAM and filtered at a 2 fold-change cut-off.

**Two pass / two-class SAM**

For this analysis two consecutive two-class SAMs were drawn: in the first pass, we searched for a “vaccination signature” by testing for differential expression in PBMCs obtained from donors after vaccination and the mature DCs included in the vaccine. Since each patient received autologous in-vitro matured DCs, we performed a paired PBMC vs matured DC two-class analysis. Only significant probes were analyzed in the second pass, consisting in an unpaired two-class (long vs short survivors) SAM. Also, in this case, we selected probes at 90th percentile FDR = 0 to statistically exclude the detection of false positive results. A 2 fold change cut-off was used for further selection.

**Identification of valid housekeeping reference genes for qPCR normalization from microarray data**

Established housekeeping genes for qPCR normalization of PBMCs samples are not consistently reported in literature. Genes that are routinely used as reference genes such as GAPDH do not perform well on whole blood or PBMC samples [65, 66]. We therefore employed our microarray dataset to extract housekeeping genes to be used as a reference for qPCR. We first scanned the expression values of genes available in the microarray platform across all PBMC samples, searching for invariance (standard deviation < 5 %) in all genes with a reasonable deviation from background expression level, to avoid the selection of non-expressed genes. Best-performing genes were then further screened for absence of long/short survivors trends by analyzing both the significance of T-tests and Mann-Whitney tests, the slopes associated to a linear regression of long versus short survivor samples and Mann-Whitney tests, the slopes associated to a linear regression of long versus short survivor samples and maximal overlap between the long and short survivor samples. Via this method we isolated PBGD, EEF1A1 and OXSR1 genes as most stably expressed genes. After measuring by qPCR the expression levels of these MA-derived three genes plus the frequently reported GAPDH gene in samples already used in microarrays, we used the procedure described by Vandesoepele and coworkers that relies on pairwise variations across samples to rank HKs genes according to their stability and perform a progressive, stepwise elimination [67]. Following this selection, two reference genes were validated, namely OXSR1 and PBGD, and used in combination as reference genes for further qPCR analysis. All the statistics were calculated using the R software.

**RNA isolation and quantitative PCR on patient PBMCs**

PBMCs were obtained from each of the Center’s Biobanks. RNA was isolated using the RNeasy mini kit (Qiagen, array samples) or Trizol (Life Technologies, additional qPCR samples) following manufacturers protocol. RNA quantity was determined on NanoDrop 2000c (Thermo Scientific) and RNA quality was determined via agarose gel electrophoresis. 2 ug of RNA was DNase I treated to remove residual genomic DNA and reverse transcribed into cDNA by M-MLV reverse transcriptase (Life Technologies) to obtain 25ul of cDNA. cDNA was diluted 25x in nuclease free water. For each reaction, 4ul diluted cDNA, 300nM primers, 10ul SYBR Green (Roche) and water were added to a final volume of 20ul. Each sample was amplified using a CFX96 sequence detection system (BioRad). The following qPCR cycling conditions were used: 50°C/2min, 95°C/10min, 40cycles of 95°C/15s; 60°C/1min, melt analysis 60°C - 95°C with increment 0.5°C/5s. The gene-specific oligonucleotide primers used to determine the expression of the genes of interest are listed in supplemental table S4. To increase the chance of consistency qPCR primers were based on the MA probes with highest differential expression. PCR products were monitored by measuring the increase in fluorescence caused by binding of SYBR Green. Quantitative PCR data were analyzed using CFX96 manager and relative expression of the gene of interest was determined using the cycle threshold method (Livak KJ, 2001) with PBGD and OXSR1 as reference genes. Statistical analysis of normalized qPCR data was performed using graph pad prism software using the statistical tests indicated in the figure legends.

**Gene Set Enrichment analysis of genes correlating with PEBP1 expression, Protein-Protein Interaction, and Gene Ontology analysis**

Using the full microarray dataset, the Pearson correlation and p-value of all probes in relation to PEBP1 expression on the array was determined in the R programming environment. Genes were ranked based on their log-transformed Pearson correlation coefficient, and this ranked list was used as input for Gene Set Enrichment Analysis (GSEA) against the blood transcriptional modules described by LI at al. [35–37]. Alternatively a ranked list of the correlation of human genes with PEBP1 in the immune-navigator database (http://sysimm.ifrec.osaka-u.ac.jp/immuno-navigator/) was used as input. For GSEA standard operating setting was applied taking into account only the highest ranking gene sets with a minimum 10 genes.

To build PPI networks the top 500 of genes, correlating or anti-correlating with PEBP1 expression were loaded into STRING database (http://string-db.
(org) to find any direct (physical) or indirect (functional) associations [68]. Using this database we simultaneously mapped genes on gene-ontologies. PPIs with a high confidence score were loaded in Biolayout express(3) for visualization (http://www.biolayout.org/) [69].

**Abbreviations**

delayed type hypersensitivity response (DTH), Dendritic cell (DC), microarray (MA), myeloid-derived suppressor cells (MDSCs), peripheral blood mononuclear cells (PBMCs), phosphatidylethanolamine binding protein 1 (PEBP1), Raf Kinase inhibitory protein (RKIP), receiver-operator curves (ROC), statistical analysis of microarray (SAM)

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**CONFLICTS OF INTEREST**

The authors declare no conflicts of interest

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