

ARTICLE

Impact of checkpoint blockade on cancer vaccine-activated CD8⁺ T cell responses

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Immune and molecular profiling of CD8 T cells of patients receiving DC vaccines expressing three full-length melanoma antigens (MAs) was performed. Antigen expression levels in DCs had no significant impact on T cell or clinical responses. Patients who received checkpoint blockade before DC vaccination had higher baseline MA-specific CD8 T cell responses but no evidence for improved functional responses to the vaccine. Patients who showed the best clinical responses had low PD-1 expression on MA-specific T cells before and after DC vaccination; however, blockade of PD-1 during antigen presentation by DC had minimal functional impact on PD-1^{high} MA-specific T cells. Gene and protein expression analyses in lymphocytes and tumor samples identified critical immunoregulatory pathways, including CTLA-4 and PD-1. High immune checkpoint gene expression networks correlated with inferior clinical outcomes. Soluble serum PD-L2 showed suggestive positive association with improved outcome. These findings show that checkpoint molecular pathways are critical for vaccine outcomes and suggest specific sequencing of vaccine combinations.

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Introduction

Melanoma represents <1% of skin cancers but is the most lethal. Over 96,000 new cases of melanoma were expected in 2019, a continued increase in incidence for the past 30 yr (Siegel et al., 2019). Checkpoint blockade therapy using monoclonal antibodies against CTLA-4 or PD-1/PD-L1 has shown durable responses as monotherapy, and as CTLA-4⁺PD-1 combination blockade, responses are seen in more than half of patients treated (Larkin, et al., 2015). However, there is still a significant proportion of patients who do not respond to checkpoint blockade or acquire resistance to these therapies (Sharma et al., 2017). Therefore, the need to develop and promote effective antitumor immunity and rationally design combination approaches is a critical issue.

The aim of dendritic cell (DC) vaccination is to induce or expand functional and long-lived tumor-specific T cell responses. Numerous studies have shown the safety and immunogenicity of DC vaccines in melanoma and other cancers (Santos and Butterfield, 2018; Gross et al., 2017; van der Burg et al., 2016; Carreno et al., 2015; Palucka and Banchereau, 2013). Despite these findings, objective clinical response rates remain low at <15% (Anguille et al., 2014). Thus, there is a need to optimize vaccine design to improve clinical responses by

promoting more effective long-term antitumor immunity and to define where vaccination could be fruitfully combined with complementary approaches such as checkpoint blockade.

In this study, CD8 T cells specific for melanoma antigens (MAs) were analyzed from patients who received autologous DCs engineered to express three full-length MAs (tyrosinase, MART-1, and MAGE-A6; Butterfield et al., 2019). The immunological and clinical impacts of key checkpoint molecule expression by the T cells were examined. Differential expression analyses in circulating lymphocytes and tumor samples, performed using a defined subset of cancer and immune-related gene and protein panels, suggest important molecular pathways for modulation and optimal sequencing of future combination trials with vaccines.

Results

Antigen expression level in the DC vaccines has minimal impact

To test the hypothesis that antigen expression levels are important for induction of T cell responses, the vaccines were

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engineered by a single adenovirus (AdV) to express high levels of tyrosinase, intermediate levels of MART-1, and low levels of MAGE-A6. Tyrosinase and MART-1 mRNAs were driven by the CMV promoter, and their expression was significantly correlated ($P = 0.00013$). MAGE-A6 mRNA was driven by the Rous sarcoma virus promoter and did not show a statistically significant correlation with the other two antigens (Fig. S1 A). Each patient's vaccine cells were transduced with differing efficiency; hence, there were individualized levels of each antigen. Cox regression analysis did not reveal significant associations between antigen expression dose in the vaccines and overall survival (OS) or progression-free survival (PFS; Fig. S1 B).

MA-specific CD8 and CD4 T cell responses

To confirm that the DC vaccines induced and/or activated tumor-specific CTL responses, ex vivo ELISPOT assays from total peripheral blood mononuclear cell (PBMC) cultures demonstrated that 20/31, 19/31, and 14/31 patients showed increases in IFN- γ -producing T cells in response to tyrosinase, MART-1, or MAGE-A6, respectively, indicating successful vaccination (Fig. 1 A, left panel). Furthermore, we show that about half of the patients had preexisting functional T cell responses to MAs before DC vaccination (Fig. 1 A, right panel). There were no significant associations between antigen-specific T cell responses and OS or PFS (Fig. S1 C).

16 patients were HLA-A2 $^+$; hence, HLA-A*0201 dextramers were used to identify peptide-specific CD8 T cell responses to Tyros_{368–376}, MAGE-A6_{271–279}, and MART1_{27–35} (Fig. 1 B). As with the trend observed with ELISPOT data, the majority of the 16 patients had dextramer-positive CD8 T cells in the peripheral blood at baseline (Fig. 1 C). As a positive control, nearly all (15/16) patients exhibited baseline reactivity to FluM1_{58–66}. Overall, three patients showed increases to three of three peptides, six to two of three peptides, and five to one of three peptides, and two (both progressive disease [PD]) had no frequency increases. Testing CD4 T cells for reactivity to full-length antigens showed a transient increase in postvaccine MA-specific activation of CD4 T cells; MART-1 and AdV-specific responses had the strongest evidence for a difference between baseline and day 43 (d43) measurements (Fig. 2 A). We did not observe any consistent trend in patient CD4 T cell tumor antigen reactivity among the response groups after vaccination. Furthermore, there were no statistically significant correlations between vaccine-induced tumor antigen-specific CD4 and CD8 T cell responses (Fig. 2 B).

The importance of vaccine antigen-specific responses has been unclear across trials due in part to patient and assay heterogeneity. As shown above, individual antigen responses did not significantly impact OS or PFS. In contrast, when bulk PBMC, purified CD8 or CD4 T cells, or CD8+CD4 T cell responses to all three shared antigens were combined, significant associations with survival could be observed (Fig. 3, A and B). Kaplan-Meier curves comparing patients whose T cells did or did not exhibit antigen-specific responses (to any of the MAs) showed that having a CD8 T cell response was associated with longer OS and PFS, and this association was also apparent when considering the combined CD8 and CD4 T cell responses (Fig. 3 B). This provides evidence that total vaccine-induced T cell responses are important in determining patient outcome.

Because this trial allowed enrollment of late-stage disease, 26/35 patients enrolled in the trial had received one or more prior systemic treatments. 15 patients had received CTLA-4 checkpoint blockade, seven of whom also received PD-1 blockade (Table S1; Butterfield et al., 2019). ELISPOT counts and dextramer-positive T cell frequencies used to measure MA-specific T cell responses at multiple time points (Fig. 4) appeared to be higher overall in patients who had received previous α -CTLA-4 and/or α -PD-1 therapies (Table S2). To evaluate this effect along with changes in MA-specific T cell responses over time after vaccination, we used a linear mixed effect model with fixed effects for time and previous checkpoint inhibitor therapy and random intercepts per patient to account for correlation of measurements within individuals. For these 16 HLA-A2 $^+$ patients, ELISPOT counts representing IFN- γ response were higher by \sim 6 per 10⁵ T cells in patients who had received previous checkpoint inhibitor therapy, and the dextramer frequencies showed a small but significant difference based on previous therapy for MART1_{27–35}. These data suggest that blockade of CTLA-4 and/or PD-1 can slightly amplify MA-specific T cells, and this difference can persist through the vaccine time course (day 0–89 [d0–d89]). However, these responses are not amplified by subsequent vaccination. The only antigen-specific T cell response that showed a clear increase over time was the MAGE-A6_{271–279} dextramer frequency, but not its corresponding functional ELISPOT responses.

Patients with tumor regression have vaccine antigen-specific CD8 T cells with low levels of PD-1 expression

To address the role of activation and exhaustion phenotypes on MA-specific CD8 T cells, surface expression of PD-1, CTLA-4, TIM-3, and LAG-3 was tested (Fig. S2 A). Two partial responders (PRs; patients 10 and 20) had strikingly low frequencies of PD-1-positive total and MA-specific CD8 T cell populations (Fig. 5 A, B). Patient 34 with stable disease (SD) also exhibited low PD-1 expression. Patient 21, a patient who had no evidence of disease (NED) at enrollment and who then progressed at 7 mo, also exhibited low PD-1 expression at baseline but showed 25%–41% positive PD-1 expression by d89. Only patient 34 had received α -PD-1 therapy (the other three patients were treatment naive or had previous CTLA-4 blockade or previous IFN- α). Expression of CTLA-4 (Fig. 5 C) was less frequent than PD-1, and there were no significant correlations observed between clinical outcome and expression of CTLA-4 and/or TIM3 on MA-specific CD8 T cells (TIM3 data not shown). Because PD-1 is also an activation marker, the IFN- γ ELISPOT results for these four patients were examined to evaluate whether there was evidence of MA-specific T cell activation. Three of four had CD8 T cell responses to one to three MAs, but at just over the limit of detection, while the fourth had a total PBMC (CD4 or CD8) response. Thus, the lack of PD-1 protein detection on total and MA-specific CD8 T cells may be due to low activated MA T cell frequency.

Long-term follow-up of melanoma-specific T cell responses in two patients

Of the 35 patients, 14 (40%) were alive at follow-up assessment. Post-trial PBMC samples were available for HLA-A2 $^+$ patients 10 and 18 from 1.5 and 2 yr after DC vaccination, respectively, which were tested for long-term MA-specific CD8 T cell

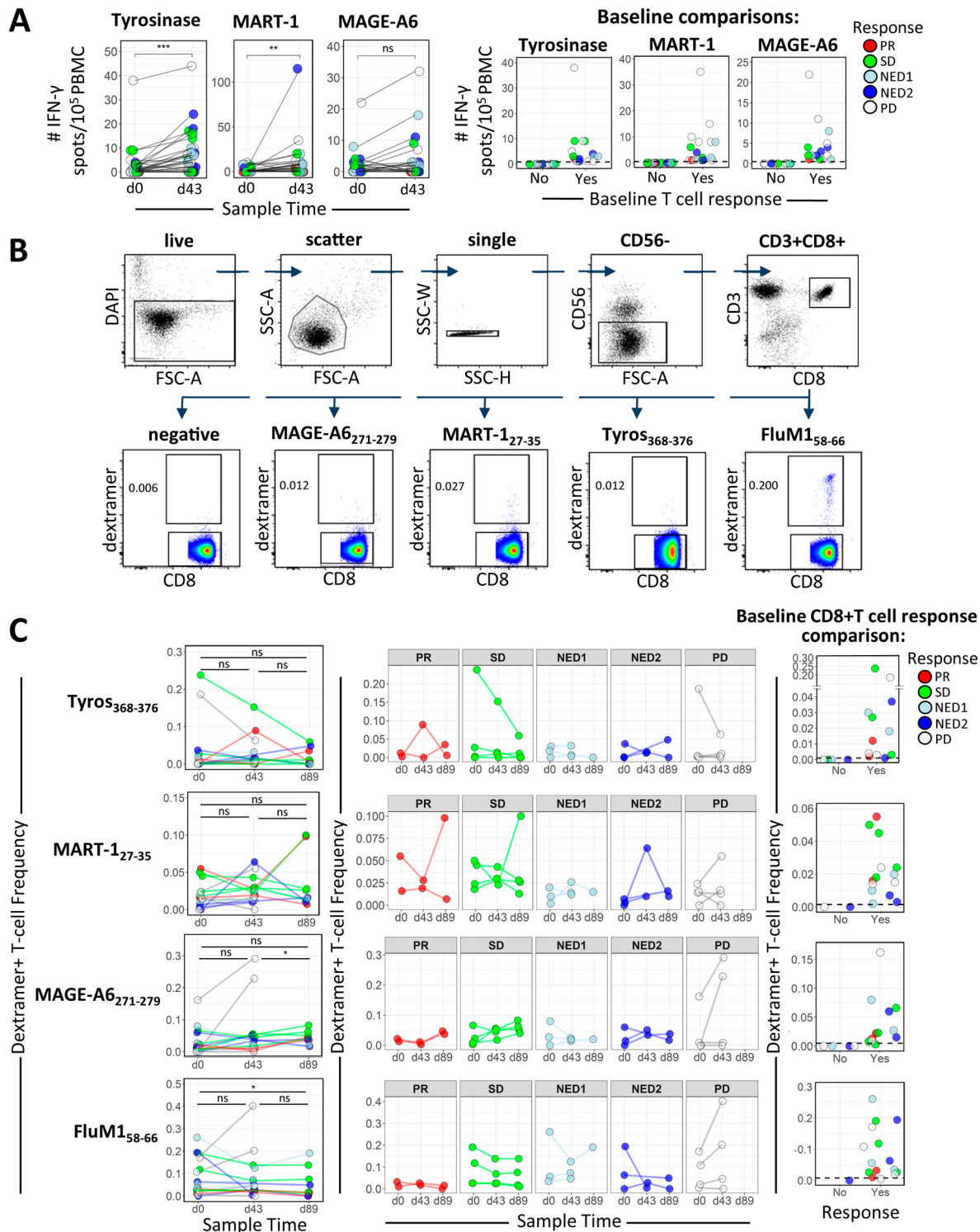
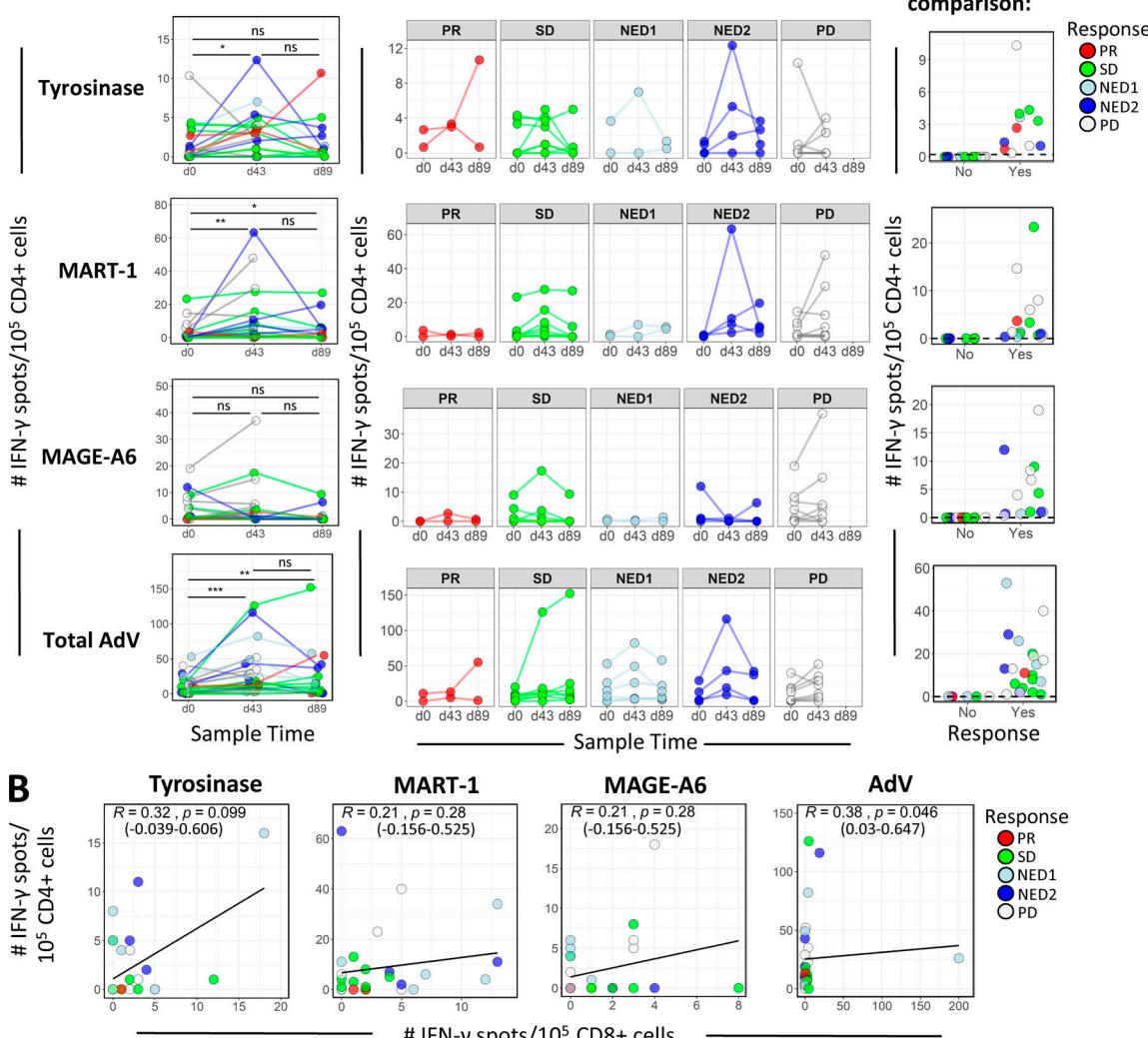


Figure 1. MA-specific responses before and after DC vaccination. **(A)** In the left panel, PBMCs from patients before (d0) and after (d43) DC vaccination were tested by ELISPOT assay for IFN- γ -producing T cells with specificities against the indicated MAs encoded in the DC vaccine ($n = 31$). Changes in influenza (FluM1) peptide-specific T cells are shown for comparison. P values were calculated using the Wilcoxon signed rank test. The right panel shows baseline IFN- γ ELISPOT count distributions separated based on the absence (No) or presence (Yes) of positive responses to individual MA antigens. **(B and C)** Gating strategy (B) and summary graphs showing the frequency of dextramer-specific CD8 T cells in the peripheral blood of HLA-A2 $^+$ patients (C) measured at d0, d43, and d89 as indicated ($n = 16$). P values for comparisons of frequency distributions between time points were calculated using Wilcoxon signed rank tests. **(C)** The middle panel shows dextramer-specific CD8 frequencies grouped according to clinical response. Baseline frequency distributions based on the absence (No) or presence (Yes) of positive responses to individual MA antigens are shown in the right panel. Positive responses were detected for 12/16, 12/16, 14/16, and 15/16 patients for Tyros₃₆₈₋₃₇₆, MART1₂₇₋₃₅, MAGE-A6₂₇₁₋₂₇₉, and FluM1₅₈₋₆₆, respectively. *, $P \leq 0.05$; **, $P \leq 0.01$; ***, $P \leq 0.001$. ns, not significant; Tyros, tyrosinase; FSC-A, forward scatter area; SSC-H, side scatter height; SSC-W, side scatter width; SSC-A, side scatter area.

A



B

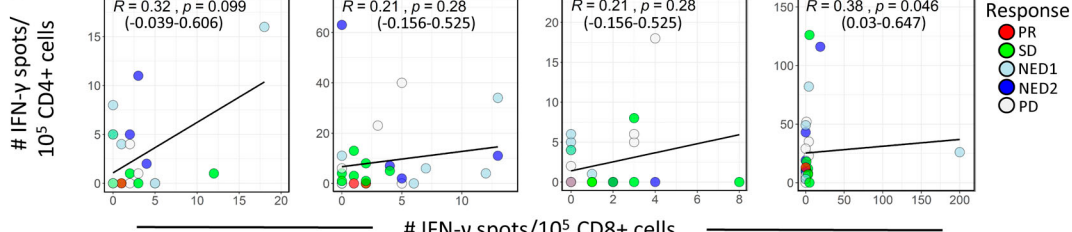


Figure 2. CD4 T cell IFN- γ ELISPOT frequencies and their correlations with CD8 responses. (A) CD4 cells from patients at d0, d43, and d89 were tested by ELISPOT assay for IFN- γ responses specific against the indicated MAs encoded in the DC vaccine ($n = 35$). P values for comparisons of frequency distributions between time points were calculated using Wilcoxon signed rank tests. The middle panel shows antigen-specific CD4 IFN- γ ELISPOT counts grouped according to clinical response. Baseline frequency distributions based on the absence (No) or presence (Yes) of positive responses to individual MA antigens are shown in the right panel. Positive responses were detected for 11/24, 15/24, 13/24, and 18/24 patients for tyrosinase, MART-1, MAGE-A6, and total AdV, respectively. *, $P \leq 0.05$; **, $P \leq 0.01$; ***, $P \leq 0.001$. (B) Scatter plots showing correlations between CD4 and CD8 after vaccine (d43); IFN- γ ELISPOT counts reactive to full-length antigens: tyrosinase (equivalence test for $R > -0.3$ and $R < 0.3$; $P = 0.547$), MART-1 (equivalence test for $R > -0.3$ and $R < 0.3$; $P = 0.305$), MAGE-A6 (equivalence test for $R > -0.3$ and $R < 0.3$; $P = 0.305$), and total AdV (equivalence test for $R > -0.3$ and $R < 0.3$; $P = 0.215$). Spearman correlation coefficient (R), P values based on asymptotic t approximation, and 95% confidence intervals are indicated. ns, not significant.

responses (Fig. S2 A). Patient 10 (partial response [PR]) subsequently received pembrolizumab for 5 mo. At 1.5 yr after vaccination, this patient had 166- and 40-fold increases in the frequency of MAGE and MART-1 circulating CD8 T cells (0.50% MAGE-A6_{271–279}⁺ and 1.00% MART1₂₇–35⁺) compared with d43. There was also higher PD-1 expression on total CD8 and MA-specific T cells (0%–5% vs. 22%–26%). LAG3 was minimal (Fig. S2 A). 4.5 yr after DC vaccination, this patient has multiple soft tissue and muscle metastases.

Patient 18 had NED on initial assessment following the protocol intervention, then progressed within 5 mo and received ipilimumab for 3 mo followed by pembrolizumab. 2 yr after vaccination, this patient had 100-fold higher

MART1_{27–35}⁺ CD8 T cells in the periphery. There was 13%–16% PD-1 expression on CD8 and MART1_{27–35}-specific T cells (Fig. S2 A), similar to d43 (13.5%–17% PD-1⁺). There was also minimal LAG3 expression detected. At 4.75 yr from vaccination, this patient has measurable disease and is now in a different clinical trial. These data suggest that postvaccine checkpoint blockade, particularly anti-PD-1, may amplify vaccine-induced T cell responses.

Simultaneous PD-1 blockade and MART-1 antigen expression by DCs

To determine whether enforced PD-1 reduction (Shimizu et al., 2018) at the same time as antigen presentation would enhance

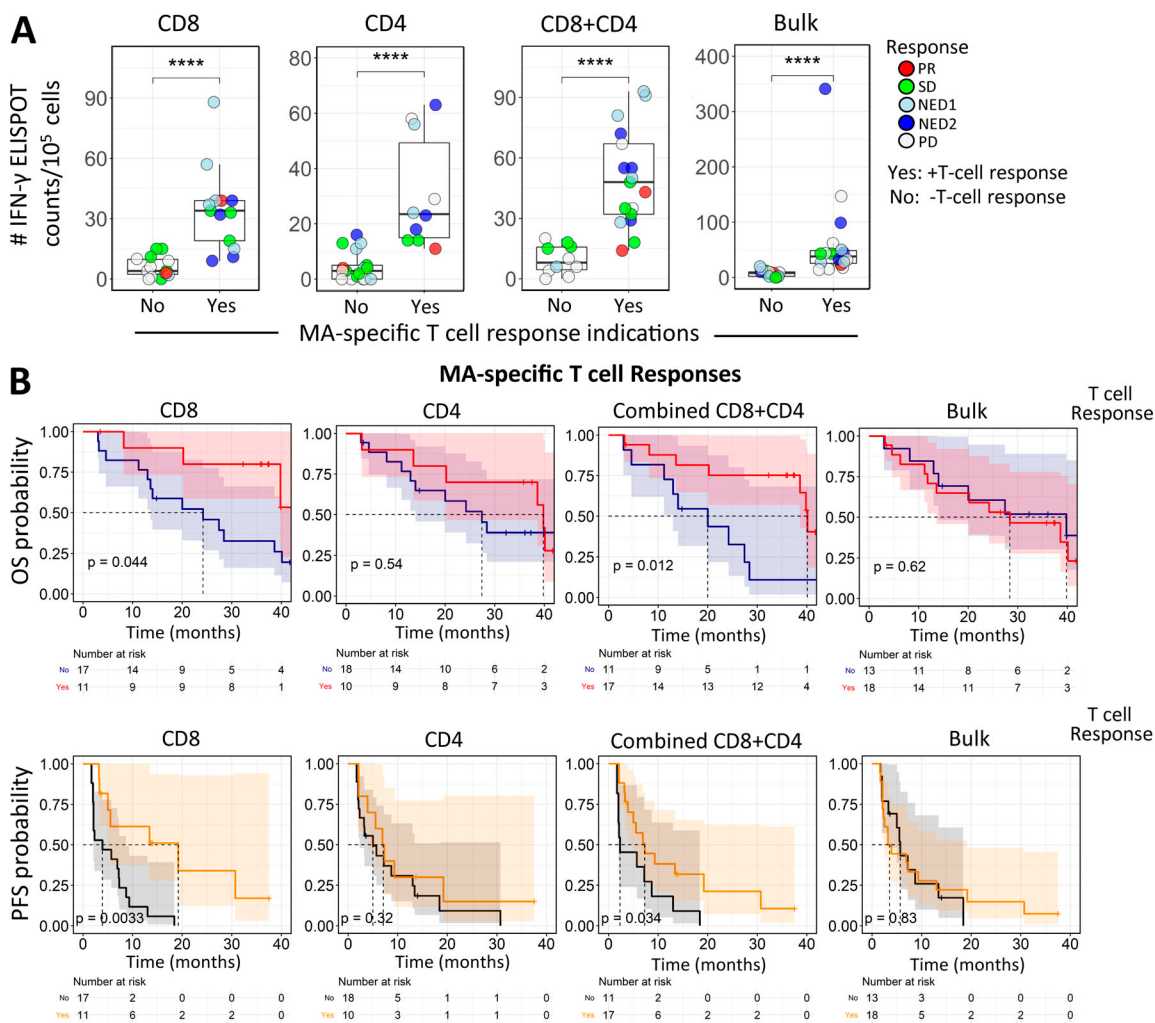


Figure 3. Combined totals of individual antigen-specific T cell IFN- γ ELISPOT counts correlate with PFS. (A) Summary graphs for CD8, CD4, total (CD8+CD4), and bulk (total PBMCs) IFN- γ ELISPOT counts (net count total for the three shared antigens at d43) with respect to T cell responses, as defined in Materials and methods. Test groups: CD8⁺ selected T cells responding to either autologous “iDCs” (immature DCs) transduced with a single MA-encoding AdV or (in HLA-A2⁺ patients) T2 cells pulsed with well-characterized immunodominant peptides; CD4⁺ selected T cells responding to autologous iDCs transduced with single MA AdV; and bulk (total PBMCs) T cell responses to autologous iDCs transduced with single MA AdV. ****, P ≤ 0.0001. (B) Kaplan-Meier survival analysis of OS and PFS comparing the survival benefits of positive CD8, CD4, combined CD8+CD4, and bulk responses. The log rank test was used to compare the Kaplan-Meier curves.

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T cell responses, patients' T cells with high PD-1 expression were stimulated in vitro with autologous DCs transduced with AdV-MART1 (Butterfield et al., 1998), then restimulated with DCs transduced with AdVMART1 ± Ad5.hPD1Ab (encoding a secreted form of human α-PD-1 with an IgG4 isotype (Garcia-Bates et al., 2016; Fig. S2 B). There were no significant differences in CD8 T cell proliferation or in expansion of MART1_{27–35}⁺ CD8 T cells (data not shown). T cells stimulated with DCs transduced with AdVMART1 + Ad5.hPD1Ab had reduced PD-1⁺-expressing CD8 and MART1_{27–35}⁺ T cells by frequency and mean fluorescence intensity, suggesting successful PD-1 blockade (Fig. 6, A and B). MART1_{27–35}⁺ T cells stimulated with DCs transduced with both adenoviral constructs also had significantly more LAG3⁺ as well as LAG3⁺TIM3⁺-expressing cells (Fig. 6 B). There were no significant differences in IFN- γ or TNF- α production (Fig. 6 C). T cells that were stimulated with both constructs showed

reduced frequency of CD69⁺ CD8 T cells, suggesting decreased activation upon addition of α-PD-1.

Protein profiling of circulating lymphocytes

To more broadly examine key immune mechanisms, targeted protein levels in circulating lymphocytes were analyzed using a 30-plex NanoString panel, and differential protein expression was compared between clinical outcomes and functional T cell responses. Table 1 depicts differentially expressed proteins between outcome groups in both baseline and postvaccine samples. With a focus on checkpoint expression, we observed significantly higher PD-1 protein counts in the unfavorable (bad) patient outcome group at d43 (Fig. 7 A). CTLA-4 levels showed similar elevation in the bad outcome groups for both d0 and d43, although these comparisons were not significant at $\alpha = 0.05$ (d0 mean count difference = 316; d43 mean count difference = 117;

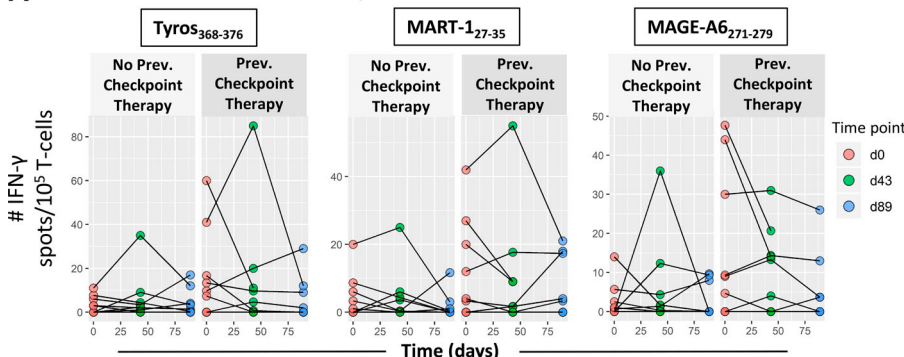
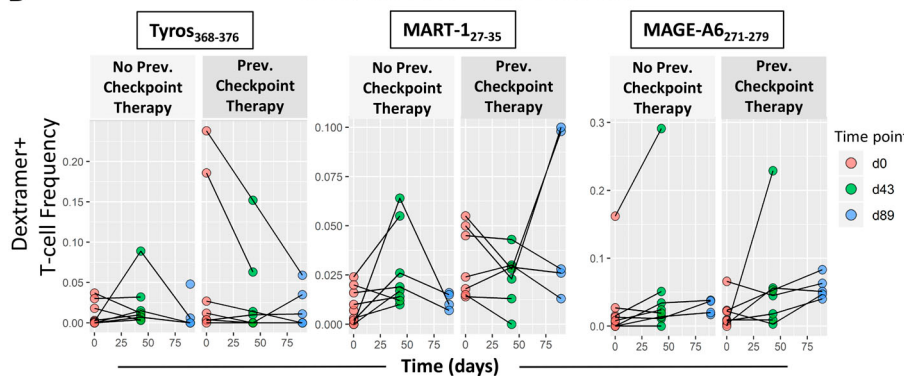
A**MA-specific CD8+ T cell ELISPOT****B****MA-specific CD8+ T cell Dextramer**

Figure 4. Effect of previous treatment with checkpoint blockade on MA-specific T cell responses. Melanoma patients were plotted separately based on whether patients received checkpoint blockade before trial enrollment and DC vaccination. Patients in the “Previous Checkpoint Therapy” panels received α -CTLA-4 and/or α -PD-1 before DC vaccination. **(A)** IFN- γ ELISPOT counts for CD8 T cell responses against Tyros₃₆₈₋₃₇₆, MAGE-A6₂₇₁₋₂₇₉, and MART-1₂₇₋₃₅ ($n = 28$; 15 no blockade, 13 with blockade) were compared between patient groups across the entire time course. **(B)** Frequencies of dextramer-specific CD8 T cells for Tyros₃₆₈₋₃₇₆, MAGE-A6₂₇₁₋₂₇₉, and MART-1₂₇₋₃₅ ($n = 16$; 9 no blockade, 7 with blockade) were compared between patient groups across the entire time course. **(A and B)** Differences were characterized by linear mixed model analysis, with results listed in Table 1. Prev., previous; Tyros, tyrosinase.

Fig. 7 A). Comparing checkpoint expression levels from all time points across the PD, SD, and PR outcome groups shows incrementally higher checkpoint expression in worse outcomes (PR < SD < PD; **Fig. S2 C**). To further evaluate the differential checkpoint expression between outcome groups over the entire time course (**Fig. 7 B**), we used a linear mixed effect model with fixed effects for time and outcome and random intercepts per patient to account for correlation of measurements within individuals. The outcome terms in these models did not show $P < 0.05$ by a likelihood ratio test, but the data nevertheless appear consistent with higher expression levels in the unfavorable outcome group (**Fig. 7 B**). Furthermore, Cox regression analysis revealed significant inverse associations between baseline CTLA-4 expression and OS and PFS (**Fig. 7 C**). In postvaccine measurements, higher CTLA-4 levels were associated with lower OS (hazard ratio [HR] of 2.2 based on standardized expression levels; $P = 0.03$), while higher expression of PD-1 appeared to be associated with worse PFS (HR of 2; $P = 0.06$; **Fig. 7 C**). Next, we compared checkpoint expression based on patient-derived MA-specific T cell responses. Significantly higher CTLA-4 protein expression was observed for patients who showed an IFN- γ response in bulk PBMCs, but not in purified T cell subsets (**Fig. S3 A**). We did not observe significant differences in PD-1 levels for bulk PBMC or purified T cell subsets.

Analysis of soluble cytokines revealed that patients in more favorable outcome groups (PR/SD) exhibited higher levels of soluble PD-L2 (sPD-L2) and lower levels of the exhaustion factor sLAG3 (**Fig. 7 E** and **Fig. S3 B**). Expression of sPD-L2 was inversely correlated with sLAG3 and positively correlated with sTIM3 (**Fig. 7 E**). A Cox regression model incorporating time-

dependent covariates was used to evaluate the effect of sPD-L2, as well as sPD-1, sTIM3, and sLAG3, on patient survival. When these soluble proteins were tested in univariate models, sPD-L2 showed the best evidence for association with OS (HR of 0.65; $P = 0.08$), and higher levels were associated with longer survival (Table S5).

Gene expression analysis of circulating lymphocytes and tumors

Gene expression analysis of circulating lymphocytes at baseline and d43 revealed 11 and 63 significantly differentially expressed genes, respectively (**Fig. 8, A and B**; and **Fig. S4 A**). Differentially expressed genes in the favorable response group at baseline were enriched for immunological pathways pertaining to TCR signaling, chemokine signaling, tumor rejection, and immune memory (**Fig. 8 A**). Favorable response groups in d43 samples showed enrichment for B cell receptor, adaptive immunity, and class II MHC antigen presentation signaling pathways. Of interest, a cluster of genes in the CTLA-4 inhibitory signaling and CD28 costimulatory pathways were enriched in the unfavorable response group (**Fig. 8 B**). Unexpectedly, gene sets representing type I IFN responses correlated with inferior clinical response at both baseline and d43 time points (**Fig. 8, A and B**).

In addition to differences between outcome groups, we examined immune checkpoint pathway changes between vaccine trial sample time points at d43 versus d0 and d89 versus d43. For these analyses, we used circulating lymphocytes from patients with good clinical outcome, as there were no available samples from d89 for poor outcome group patients. Here, we show that the CTLA-4 inhibitory signaling pathway was significantly

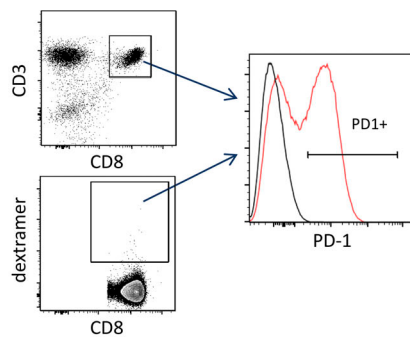
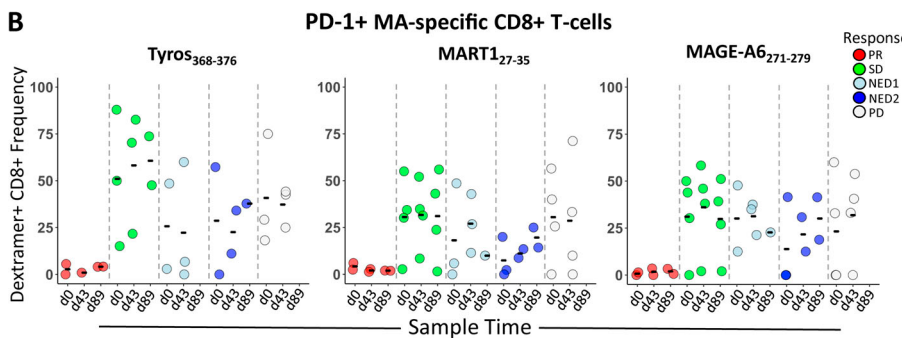
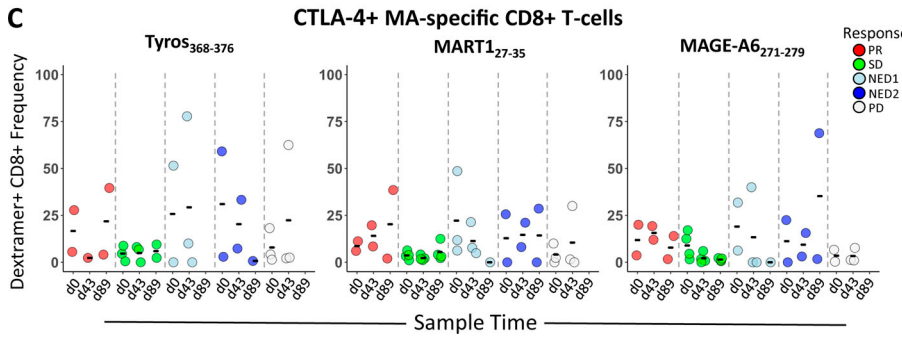
A

Figure 5. PD-1 and CTLA-4 expression on CD8 and MA-specific T cells. (A) Gating strategy used to determine dextramer frequencies of PD-1- and CTLA-4-positive MA-specific CD8 T cells from HLA-A2⁺ patients. **(B and C)** Shown are distributions of PD-1- (B) and CTLA-4-positive (C) dextramer-specific CD8 T cells from HLA-A2⁺ patients grouped according to clinical outcome across time points (d0, d43, and d89; $n = 16$ total; two PR, three SD, three NED1, three NED2, and five PD). Tyros, tyrosinase.

B**C**

down-regulated following vaccination (Fig. 8 C). We also surveyed checkpoint pathway changes between d43 and d89 and saw that the PD-1 signaling pathway was primarily down-regulated during this later part of the trial (Fig. 8 D). The analogous pathway results for PD-1 and CTLA-4 are shown in Fig. S4, B and C, respectively. With a focus on immune checkpoint signaling, we used the linear combination method within the gene set enrichment analysis (GSEA) software to determine pathway enrichment based on correlation with PD-1 or CTLA-4 protein expression as a continuous phenotype. With an interest in postvaccine-induced changes, we combined the d43 and d89 (omitting baseline samples) circulating lymphocyte expression sets for this analysis (Fig. 9 A and Fig. S4 D). Interestingly, B cell receptor pathways negatively correlated with elevated PD-1 protein expression, while gene signatures downstream of T cell receptor activation and IL2 signaling showed negative correlations with high CTLA-4 protein expression (Fig. 9 A). Furthermore, increased CTLA-4 protein levels showed strong associations with numerous IFN type I response pathways (Fig. 9 A). As a positive validation, we observed that elevated PD-1 levels correlated well with genes in the PD-1 signaling gene set (Fig. 9 A). CTLA-4 inhibitory signaling showed suggestive but not significant correlation with PD-1 protein levels (Fig. S4 D). The checkpoint pathways for PD-1 (normalized enrichment score [NES] = 0.84; P value =

0.670) and CTLA-4 (NES = 0.87; P value = 0.613) only weakly associated with the CTLA-4 protein continuous phenotype and did not show statistical significance.

In addition to circulating lymphocytes, we further evaluated clinical outcome-related gene expression changes in the CTLA-4 and PD-1 signaling pathways in patient tumor/tumor infiltrating lymphocyte (TIL) samples. Genes in the CTLA-4 and PD-1 pathways showed increased expression primarily in the inferior outcome patient groups, although this did not represent significant enrichment based on GSEA (Fig. S4 E). Additionally, we correlated checkpoint pathways in tumor samples with OS as a continuous phenotype (in months). The PD-1 pathway did not show significant enrichment in this test (NES = 0.86; P value = 0.651), but CTLA-4 inhibitory signaling in the tumor/TIL samples was negatively associated with OS (Fig. 9 B). There was no enrichment found based on correlation with OS for the PD-1 (NES = 0.59; P value = 0.952) or CTLA-4 (NES = -0.95; P value = 0.534) pathways in circulating lymphocytes.

Discussion

Here, we have tested multiple key aspects of the T cell response to cancer vaccination to identify the critical parameters for

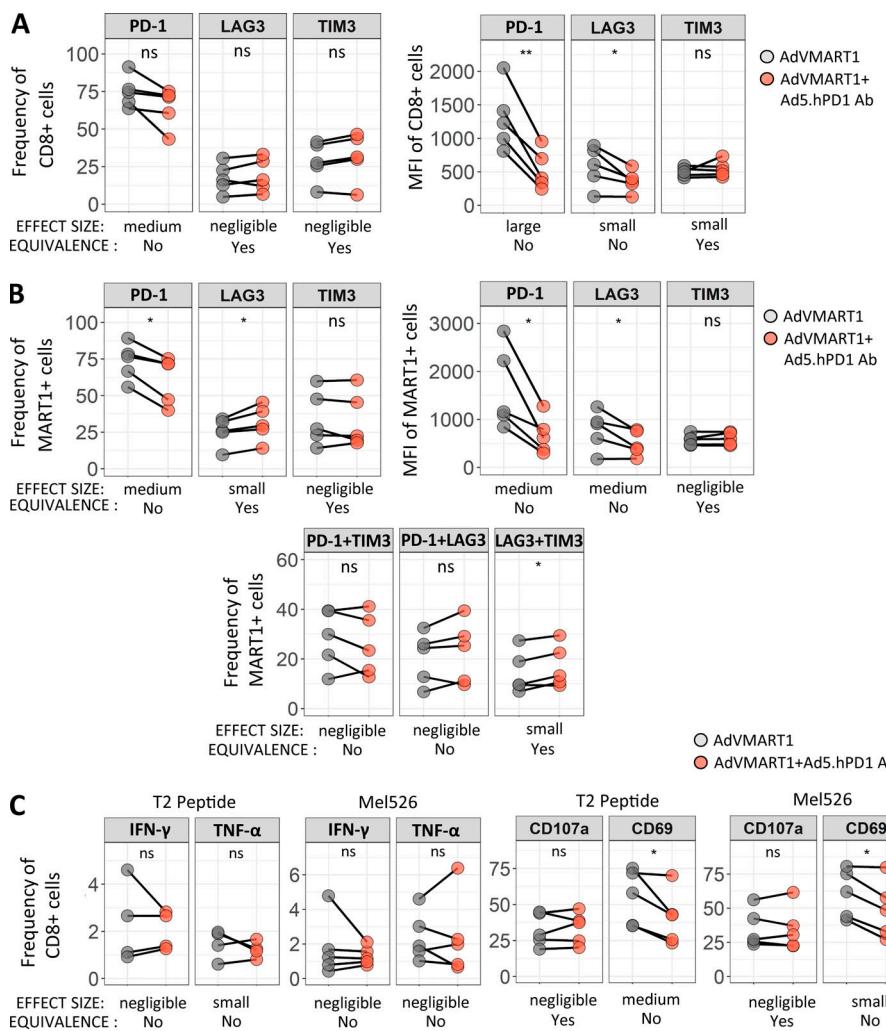


Figure 6. Effect of stimulating CD8 T cells with DCs transduced with AdVMART1 and Ad5.hPD1Ab in vitro on T cell function. T cells were stimulated with AdVMART1 transduced autologous DCs in vitro followed by restimulation with DCs transduced with AdVMART1 only (gray circles) or in combination with Ad5.hPD1Ab (orange circles). **(A and B)** Summary graphs showing frequency and mean fluorescence intensity (MFI) of checkpoint molecules CTLA-4, LAG3, PD-1, and TIM3 in CD8 T cells (A) or MART1₂₇₋₃₅⁺ CD8 T cells (B). **(C)** T cells were stimulated with MART1₂₇₋₃₅⁺-pulsed T2 or Mel526 and examined for TNF- α , IFN- γ CD107a, and CD69 expression. $n = 5$; P values shown were calculated using paired t tests. Effect sizes are summarized using Hedges' g as follows: $g < 0.2$ (negligible), $g < 0.5$ (small), $g < 0.8$ (medium), and $g > 0.8$ (large). A detailed list of effect sizes, confidence intervals, and equivalence testing results for these data are provided in Table S2. **(A-C)** *, $P \leq 0.05$; **, $P \leq 0.01$. ns, not significant.

future success of vaccine approaches. We find that the amount of antigen expressed by the DC does not define the subsequent T cell response to that antigen or the clinical outcome of vaccinated patients. We observed reduced PD-1 on CD8 T cells in the best clinical responders, and we found that blocking PD-1 in the presence of DC T cell stimulation reduces PD-1 surface expression but does not increase antigen-specific T cell activity. These results contrast with those of Hobo et al. (Hobo et al., 2013; Van den Bergh et al., 2017), who found that reducing PD-L1/L2 by siRNA \pm IL-15 trans-presentation in DC did improve expansion of MAGE-A3-specific CD8⁺ T cells in vitro from healthy donors (HDs). One key difference may be that we tested PD-1-high T cells from cancer patients.

Previous checkpoint blockade therapy did amplify vaccine antigen T cell responses, which persisted throughout the course of vaccination. However, our functional IFN- γ ELISPOT results (Fig. 4 A) suggest that previous checkpoint blockade did not prime these patients for higher functional T cell responses. A clear increase in MAGE-A6₂₇₁₋₂₇₉ dextramer⁺ frequency over time suggests that expansion of a subset of MA-specific cells was enhanced, which is under further investigation. Two patients who received postvaccine checkpoint therapy had highly amplified MA-specific T cell frequencies, suggesting that the

schedule of vaccination and checkpoint therapy is important for optimizing T cell responses. Whether the postvaccine checkpoint therapy triggered functional memory T cell responses in these patients is also under investigation. Protein expression profiling also confirmed increased levels of the checkpoint molecules PD-1 and CTLA-4 in circulating lymphocytes from patients with worse clinical outcome (Fig. 7, A and B; and Fig. S2 C). Along with the inhibitory action on T cell activation, increased expression of these immunosuppressive molecules, primarily CTLA-4 (Wei et al., 2018), is associated with inferior clinical responses, both PFS and OS (Fig. 7 C). Despite this, differences in expression levels for PD-1 and CTLA-4 in circulating lymphocytes did not associate with IFN- γ responses from bulk PBMC or purified T cell subsets. The only significant difference found was higher CTLA-4 protein expression in the group showing an IFN- γ response in bulk PBMCs (Fig. S3 A).

While soluble PD-L1 has been used as a prognostic biomarker in melanoma patients treated with checkpoint blockade therapy (Zhou et al., 2017), the pathophysiological significance of soluble forms of checkpoint molecules including PD-1, PD-L1, and PD-L2 in serum from cancer patients is not yet fully elucidated (Zhu and Lang, 2017; Appleman et al., 2018), but it is beginning to be reported. Our multiplex serum analyses revealed that patients in

Table 1. Protein profiling in circulating lymphocytes

Protein name ^a	CD name	FC ^b	Adjusted P value ^c
Baseline PD vs. SD/PR			
CTLA-4	CD152	1.8	0.388
NCAM	CD56	1.66	0.388
CD45RO	CD45RO	1.52	0.388
d43 PD vs. SD/PR			
CTLA-4	CD152	1.39	0.786
PD-1	CD279	1.59	0.743
CD45RO	CD45RO	1.69	0.450
BTLA	CD272	-1.75	0.450
CD19	CD19	-1.8	0.743
HLA-DRA	HLA-DRA	-1.84	0.450
CD40	CD40	-2.04	0.450

NanoString protein profiling was used to test elutriated circulating lymphocytes. $n = 16$ (at baseline: 1 PR, 6 SD, 9 PD), and $n = 11$ (at d43: 1 PR, 4 SD, 6 PD). Immune checkpoint molecules are in bold.

^aCommon protein name is listed. CD, cluster of differentiation name. FC, fold change.

^bFold change is shown, where positive expression is higher in PD (Bad) compared to SD/PR (good) patient group. Negative correlations are shaded.

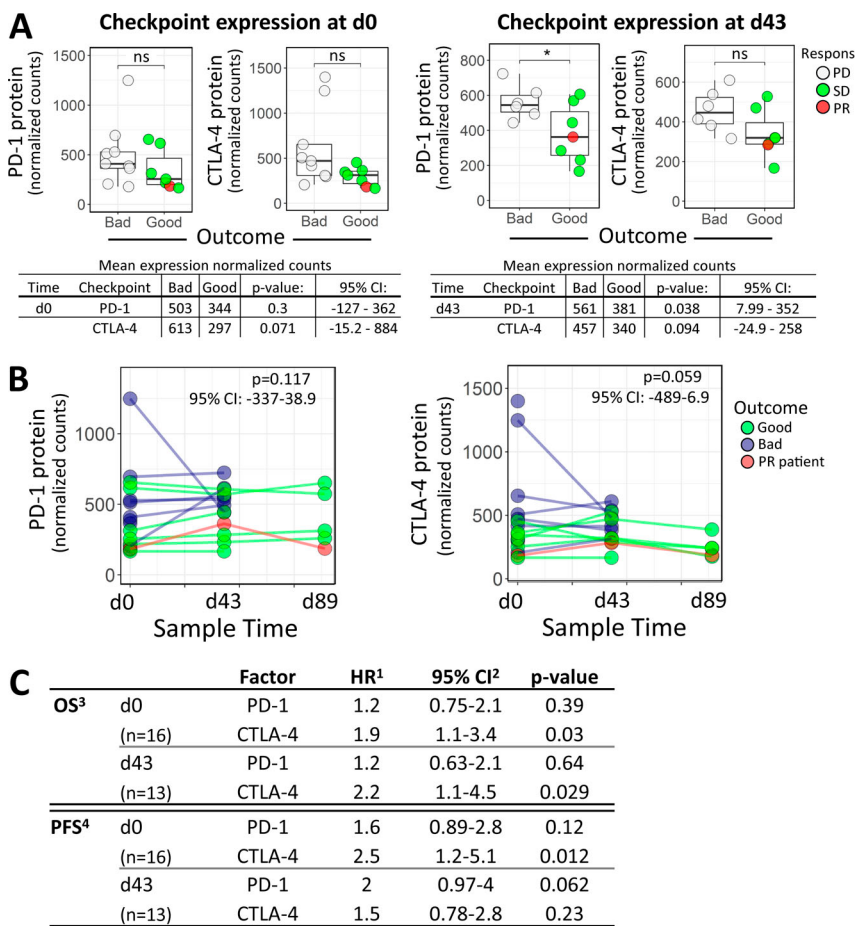
^cFalse discovery rate adjusted P value was calculated using Benjamini Hochberg correction.

more favorable outcome groups (PR/SD) exhibited increased levels of sPD-L2. Here, we report for the first time a suggestive association between higher sPD-L2 levels and longer OS in melanoma patients ($P = 0.08$; Table S5). A recent phase 2 study in hepatocellular carcinoma surveyed PD-L2 as a circulating biomarker but did not find significant correlations with clinical responses (Feun et al., 2019). We also observed that sPD-L2 correlates negatively with soluble LAG3 and positively with soluble TIM3, but the biological significance of these relationships remains to be determined (Fig. 7 E). In this study, we quantified total T cell populations by whole blood PBMC FACS, and we further showed that the amount of regulatory T cells (T reg cells) inversely correlates with sPD-L2 (Fig. S3 C). While the CD4⁺CD25⁺CD127^{low} T reg cells did not correlate with sPD-L2 levels, this T cell subtype, known to exhibit anti-proliferative effects on CD4⁺CD25⁺ T cells (Yu et al., 2012), was significantly elevated in melanoma patients compared with healthy controls at baseline and post-IFN- α treatment (Fig. S3 D). This suggests that sPD-L2 may be an important biomarker in antitumor immune responses, and it will be further explored for use in predicting vaccination efficacy in melanoma patients.

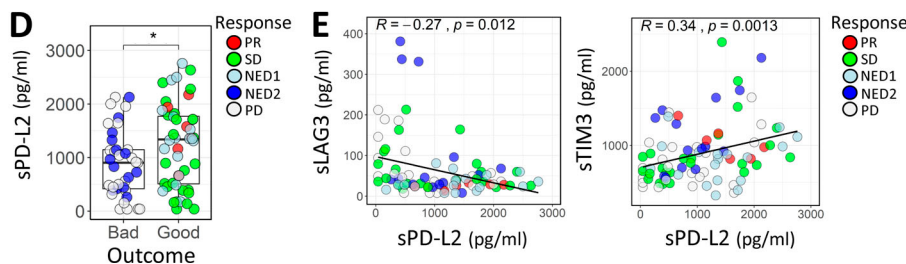
We used gene expression analysis to better define the changes associated with disease outcome and to identify possible vaccine-induced changes in circulating lymphocyte populations. Differential gene expression analysis in baseline lymphocytes showed higher expression of the IL-1 cytokine family member *IL36G* in favorable outcome patients (Fig. 8 A). *IL36G*, *CXCL19*, and *CXCL5* were part of the leading gene sets in GSEA for the IL-22 and IL-17 signaling pathways, and in a recent report by Wang et al. (2015), it was demonstrated that *IL36 γ* synergizes with IL12/TCR signaling to promote CD8 T cell function and induce antitumor immune responses. Furthermore, *IL36 γ* promoted the development of tertiary lymphoid organs and increased the

efficacy of a DC vaccine to inhibit tumor progression of melanoma tumorigenesis in vivo (Weinstein et al., 2017). Additional gene sets associated with T cell differentiation, tissue chemotaxis, and immunosurveillance and good clinical response are the IL-2 and CXCR3/4 signaling pathways (Ross and Cantrell, 2018). Chemokine receptors play a critical role in lymphocyte-mediated tumor infiltration/regression, and reduced expression of CXCR3/4 chemokine receptors on peripheral CD8 and CD4 T lymphocytes was correlated with increased metastatic properties and widespread dissemination of tumors in melanoma patients (Jacquelot et al., 2016).

In this study, we used the NanoString PanCancer Immune Profiling and chimeric antigen receptor (CAR)-T Characterization gene expression panels, which share 276 genes in common and include 478 and 494 nonshared genes, respectively. The genes *CD22*, *CD200*, *CD19*, *CD8B*, *CD40*, and *CXCR5* were shown to be up-regulated in the favorable clinical outcome groups in the postvaccine samples using both gene expression panels (Fig. S5 A). Higher *CD40* and *CD19* expression in the good response groups was also found at the protein expression level (Table 1). In addition to T cell receptor and costimulatory molecules, elevated expression of *CD22*, *CD19*, and *CD40* denotes the likely presence of B cells in our circulating lymphocyte population. In fact, using GSEA pathway analysis, we found numerous B cell receptor (BCR) signaling gene signatures that were enriched in lymphocytes from patients with favorable outcome groups (Fig. 8 B). In addition, BCR signaling pathways were inversely associated with elevated PD-1 protein expression (Fig. 9 A). This result supports previous studies arguing for the importance of B cells in antigen-driven humoral immune responses against malignant melanoma (Chiaruttini et al., 2017). As part of the T cell characterization in this report, *CD8B* and *CXCR5* were shown to mark a subset of memory CD4 T cells, which specialize



¹hazard ratio, ²confidence interval, ³overall survival, ⁴progression-free survival



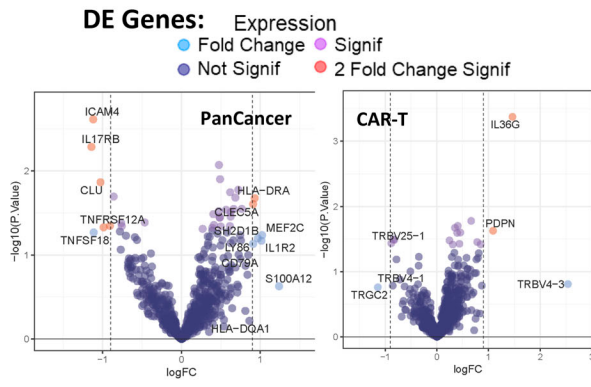
in supporting antibody-mediated immune responses (Chevalier, et al., 2011). Furthermore, a multi-parametric FACS analysis of melanoma-reactive CD4 T cells from stage III malignant melanoma patients revealed elevated CXCR5 expression in a subset of CD4 T cells with enhanced responsiveness to melanoma cell antigen reexposure (Zhang et al., 2016). In addition to its role as a receptor for T cell priming and B cell activation (Elgueta, et al., 2009), functional expression of CD40 was shown in a recent study by Levin et al. (2018) to increase IFN- γ secretion and improve the functional properties of human CD8 cells, CD4 cells, and a subset of TILs in melanoma.

In comparisons of outcome groups at specific time points, we did not detect enrichment of immune checkpoint signaling pathways using differential expression of circulating lymphocytes at baseline (Fig. 8 A). However, GSEA analyses of post-vaccine d43 samples revealed enrichment of the CTLA-4 inhibitory signaling pathway in the inferior outcome patient

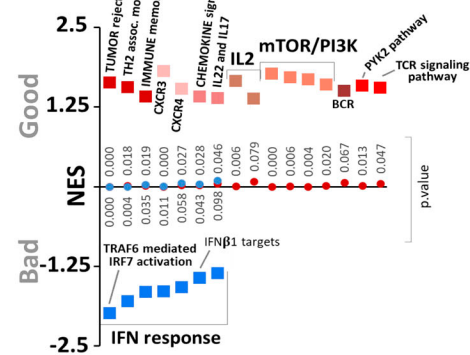
Figure 7. Immune checkpoint profiling in circulating lymphocytes and patient serum.

(A) PD-1 and CTLA-4 protein expression comparisons between outcome groups in baseline (d0) or d43 circulating lymphocytes. The Wilcoxon rank sum test and unpaired Welch's *t* test were used for calculating *P* values at d0 and d43, respectively. *, *P* ≤ 0.05. **(B)** A linear mixed effect model was used to evaluate differential PD-1 and CTLA-4 protein expression between outcome groups, accounting for repeated measures of individuals across the three time points (d0, d43, and d89). Statistical significance and 95% confidence intervals (CIs) for the outcome group coefficients are indicated. **(C)** Univariate Cox regression analysis of PD-1 and CTLA-4 checkpoint protein expression in circulating lymphocytes isolated at baseline (d0) or after vaccine (d43) and their association with OS and PFS. **(D)** Difference in sPD-L2 expression with respect to good and bad clinical outcome, with *P* value from Wilcoxon rank sum test. *, *P* ≤ 0.05. **(E)** Scatter plots showing correlations between sPD-L2 and sLAG3 (95% CI = -0.457 to -0.06) and sPD-L2 and sTIM3 (95% CI = 0.137–0.516). Spearman correlation coefficient (*R*) and corresponding *P* value are indicated on the graphs. ns, not significant.

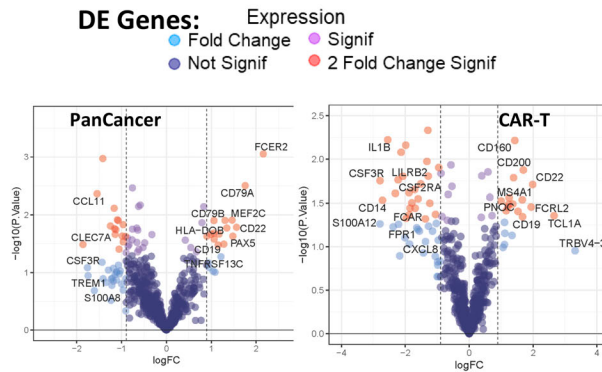
A Comparisons of SD/PR vs PD at day 0



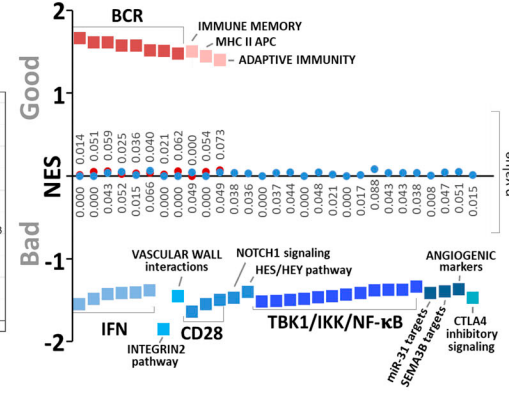
Pathway Analysis



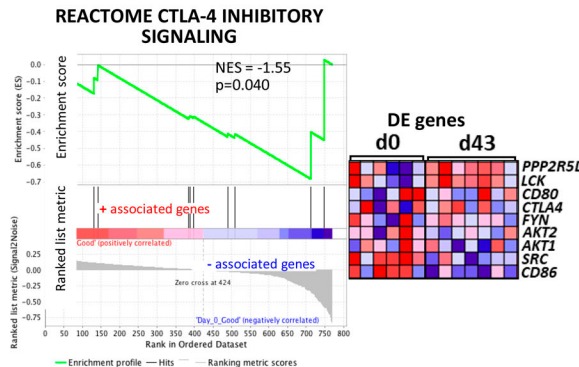
B Comparisons of SD/PR vs PD at day 43



Pathway Analysis



C d0 → d43



D d43 → d89

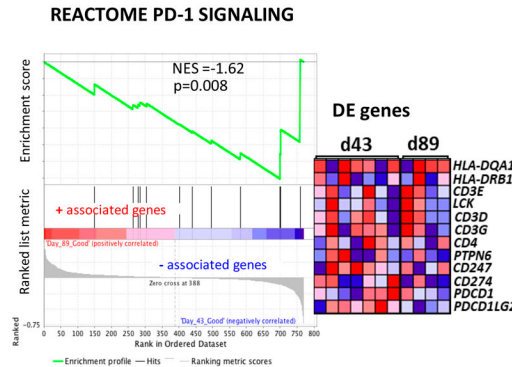


Figure 8. Differential expression and gene enrichment pathway analysis in circulating lymphocytes. (A and B) Volcano plots show differential gene expression by fold change (logFC) and P value ($-\log_{10}[p]$) for the PanCancer Immune Profiling and CAR-T Characterization NanoString gene panels comparing good (SD/PR) with bad (PD) outcome groups at baseline (d0; A) and d43 (B) in circulating lymphocyte samples. Table S3 lists information about the samples selected for NanoString profiling. Dark blue dots denote nonsignificant genes, purple dots denote significant genes with fold change ≤ 1.85 , light blue dots represent genes with fold change ≥ 1.85 and P value ≥ 0.05 , and red dots indicate genes with fold change ≥ 1.85 and P value ≤ 0.05 . Summary of pathways identified by GSEA/MSigDB analysis as showing enrichment among highly differentially expressed genes at d0 (A) and d43 (B). The y axis denotes NES, and the squares represent up-regulated (red, associated with good outcome) and down-regulated (blue, associated with bad outcome) pathways with corresponding gene enrichment P values indicated on the midline. **(C)** GSEA plot for the CTLA-4 inhibitory pathway and a heatmap of its associated gene set, illustrating enrichment for down-regulation at d43 relative to d0. **(D)** GSEA plot for the PD-1 signaling pathway and its associated gene set, which is down-regulated at d89 versus d43 in circulating lymphocytes. For C and D, circulating lymphocytes from patients with good clinical outcome were used, as there were no available samples from PD patients. Heatmaps in C and D show the clustered genes in the leading-edge subsets for each pathway category. Gene expression values are represented as colors, where the range of colors (red, pink, light blue, dark blue) shows the range of expression values (high, moderate, low, lowest).

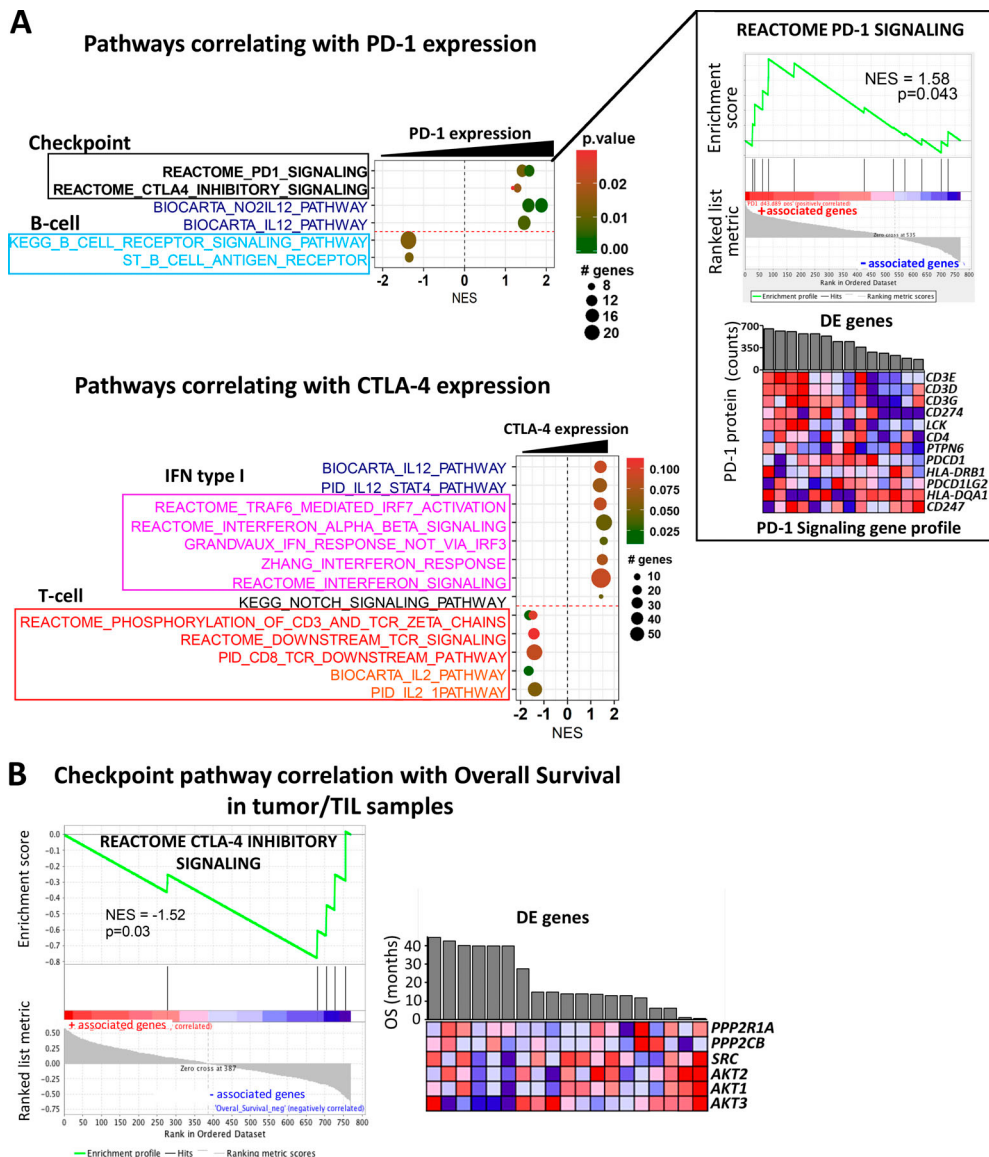


Figure 9. Significantly enriched pathways associated with immune checkpoint expression levels in circulating lymphocytes and clinical outcome in tumor biopsies. (A) Several GSEA pathways correlating with PD-1 and CTLA4 protein expression are represented. Gene sets belonging to different pathway categories are represented as dots, which are organized by their NES score along the x axis. The color and size of the dots indicate the corresponding P values and number of genes associated with the gene set, respectively. Representative GSEA plot and heatmap for genes involved in the PD-1 signaling pathway, which positively correlates with PD-1 protein expression (shown as bar graph above the heatmap) in patient lymphocytes. **(B)** GSEA plot and clustered heatmap of expression values for leading edge genes in the CTLA-4 inhibitory signaling pathway, which negatively correlates with OS in tumor biopsy samples ($n = 19$). Refer to Fig. 7 for interpretation of the gene expression values in heatmaps.

et al., 1998; Hu et al., 2001). Our gene enrichment pathway analyses and protein profiling suggest that elevated inhibitory signaling from CTLA-4 in poor outcome patients likely plays a significant role in suppressing MA-specific T cell responses to the MA-engineered DC vaccine. A combination of ipilimumab with a DC vaccine has provided encouraging and durable anti-tumor responses in a phase 2 study in pretreated advanced melanoma (Wilgenhof et al., 2016). Therefore, these data collectively suggest that a future combination approach using CTLA-4 blockade during earlier phases of vaccine administration may facilitate early stages of MA-specific T cell activation and therefore improve patient immune responses.

A clear hallmark of a poor outcome lymphocyte profile included numerous genes associated with the IFN response and TRAF6/3-mediated IRF7 activation pathways (Konno et al., 2009). Down-regulation of IFN-stimulated genes has been previously described in patients with metastatic melanoma (Critchley-Thorne et al., 2007), and importantly, changes in type I/II IFN signaling were shown to be involved in CD8 and CD4 T cell exhaustion (Ou et al., 2001; Crawford et al., 2014), which is a dysfunctional state of chronic antigen exposure in the setting of a pro-inflammatory tumor microenvironment or protracted viral infection (Baitsch et al., 2011; Wherry and Kurachi, 2015). T cell exhaustion gene signatures in the context of chronic viral

exposure were well defined in distinct subsets of CD8 and CD4 effector and memory T cell populations (Crawford et al., 2014; Wherry et al., 2007; Doering et al., 2012). We have also observed that IFN-related pathways are enriched for positive association with CTLA-4 protein expression in lymphocytes (Fig. 9 A). While the exact nature of the relationship between IFN pathway genes and bad outcomes in our trial is unclear, we hypothesize that the IFN/TRAF6-IRF7 gene sets up-regulated in the bad clinical outcome groups may be related to the molecular signatures of the T cell exhaustion phenotype. Relevant to these gene expression results, the complexity associated with type I/II IFN signaling in the context of checkpoint therapy resistance and immunosuppression in metastatic melanoma has been recently demonstrated (Benci et al., 2016; Garcia-Diaz et al., 2017). Similar to the exhaustion associated with prolonged antigen exposure, chronic IFN signaling contributes to epigenetic reprogramming of tumor cells, leading to up-regulation of inhibitory T cell ligands PD-L1 and PD-L2 on tumor and immune cells, which leads to suppression of immune surveillance and tumor progression after anti-PD-1 therapy (Benci et al., 2016; Garcia-Diaz et al., 2017). Therefore, we further surveyed our gene expression data for differences in T cell exhaustion signatures in baseline and postvaccine lymphocyte samples using the Molecular Signature Database (MSigDB) C7 immunological gene database (Fig. S5, B and C). We found that differentially expressed genes between the two outcome groups showed enrichments in several gene sets defined by contrasting exhausted CD8 T cells with other T cell subsets. The good outcome group showed enrichments for genes that are highly expressed in effector and memory CD8 T cells in contrast to exhausted T cells (Fig. S5, B-D). While it is difficult to attribute a clear exhaustion phenotype to either outcome category, it is important to ask whether the increased expression of effector/memory genes in lymphocytes may correlate with better MA responses and better clinical outcomes in patients. We also note that our analyses considered virally induced T cell exhaustion signatures, which may not fully represent changes associated with tumor-induced exhaustion and that the gene panels used here do not include a comprehensive probe set of exhaustion molecules. Future studies using FACS-based single-cell targeted technology may more precisely characterize the relationships between T cell activation and exhaustion profiles in the MA-specific T cells from distinct patient outcome groups.

In summary, our examination of lymphocyte and tumor gene and protein expression changes from baseline to 1 mo after vaccine confirmed that both immune checkpoint signaling pathways and IFN response gene networks play important roles in the response to cancer vaccines. These data, combined with the checkpoint blockade schedule effects, suggest potential future combination trials.

Materials and methods

Clinical trial and patient samples

A phase 1 single-site study was performed to evaluate the immunological effects of autologous monocyte-derived DCs transduced with tyrosinase, MART-1, and MAGE-A6 genes (by

replication-defective AdV AdVTMM2) in 35 subjects with recurrent, unresectable stage III or IV melanoma (Butterfield et al., 2019). The clinical trial reported was fully approved by the University of Pittsburgh protocol review committee and institutional review board (PRO12010416, #UPCI09-021) and had Food and Drug Administration investigational new drug #15044 and NCT01622933 (clinical data finalized April 2018). All patients and donors provided informed consent (Butterfield et al., 2019). Monocytes and lymphocytes were isolated by elutriation of the leukapheresis from each patient. The monocyte fraction was plated and differentiated into DCs with GM-CSF+IL-4 for 5 d, then matured by the addition of IFN- γ +LPS overnight. Patient vaccines were prepared by harvesting matured DCs and transducing them with AdVTMM2. Cells from baseline and after DC vaccination at d43 and after observation or 1 mo of high-dose IFN- α (d89 or 101, hereafter "89") were cryopreserved in 50% RPMI (Gibco), 40% human AB serum, and 10% DMSO (Sigma-Aldrich). HD blood samples were used as controls and obtained by venipuncture under informed consent (UPCI #04-001). HD PBMCs were purified by Ficoll gradient centrifugation and either used immediately or cryopreserved as above until use. All thawed specimens were >80% viable upon thaw. Peripheral blood flow cytometry was described previously (Butterfield et al., 2019).

Of the 35 patients who were vaccinated, 31 had ELISPOT data for PBMCs from both d0 and d43 (this testing of total PBMCs was used to check for balance in T cell responses between the observation and IFN- α trial arms); four patients progressed before collection of the d43 blood sample. 28 patients had data available from the second ELISPOT assay (testing purified CD4 $^{+}$ and CD8 $^{+}$ subsets), which was performed using samples from all three time points, but only 20 of 28 had d89 specimens due to progression (Butterfield et al., 2019). Patients were only excluded from a particular analysis if there were no specimens available or if they were not able to be tested in the assay (the MHC dextramer analysis used a reagent based on the HLA-A*0201 allele and was only performed for HLA-A2 $^{+}$ patients).

IFN- γ ELISPOT assays

MA-specific T cell responses were examined using total PBMCs at baseline and d43 to test responses to autologous DCs (10 T cells to each DC) transduced with individual antigens AdVTyrosinase, AdVMART1, AdVMAGEA6, AdVLacZ (for the AdV vector backbone response), and controls (Butterfield et al., 2019). Additional ELISPOT assays were performed on purified CD4 $^{+}$ and CD8 $^{+}$ T cells and included responses in HLA-A2 $^{+}$ patients to immunodominant peptides on T2 cells. To quantify specific responses to the melanoma-associated antigens (MA), the AdVLacZ response was subtracted from the AdV-MA response, and positivity was defined as >10 spots counted per well and at least a twofold increase over baseline.

Antibodies, dextramers, and flow cytometry analysis

Antigen-specific CD8 T cells from HLA-A2 $^{+}$ patients were identified using dextramers displaying tyrosinase₃₆₈₋₃₇₆, MAGE-A6₂₇₁₋₂₇₉, MART1₂₇₋₃₅, and influenza (FluM1₅₈₋₆₆) per manufacturer's instructions (Immudex). The lower limit of detection

was not determined, but frequencies and increases ≥ 0.003 were counted as positive. Checkpoint expression on CD8 T cells was examined using the following fluorochrome-conjugated antibodies: CD3 (UCHT1), CD8 (SK1), CD56 (NCAM16), CTLA-4 (BNI3), PD-1 (EH12; BD Biosciences), LAG3 (11C3C65; Biolegend), and TIM3 (F38-2E2; Thermo Fisher Scientific). DAPI (Molecular Probes; Invitrogen) was used as a viability dye for dead cell exclusion. FACS analyses were performed using the BD LSR Fortessa II (BD Biosciences), and data were analyzed using FlowJo v10 software.

In vitro co-cultures and intracellular cytokine assay

Monocytes were thawed and cultured for 5 d in AIMV (Gibco) supplemented with 2% human AB serum, 800 U/ml GM-CSF (Sanofi), and 500 U/ml IL4 (Gemini Bio-Products) to generate DCs. DCs were matured on day 6 with 1,000 U/ml IFN- γ (Peprotech) and 250 ng/ml LPS (Sigma-Aldrich). For the first round of stimulation, DCs were collected and transduced with AdVMART1 at 1,000 multiplicities of infection (MOI) for 3–4 h in serum-free AIMV (Butterfield et al., 1998). After transduction, DCs were washed and co-cultured with CD8 enriched T cells using magnetic separation with CD8 microbeads (Miltenyi Biotech). After 8 d, cultured T cells were collected and restimulated with a new batch of DCs transduced with AdVMART1 at 500 MOI with or without Ad5.hPD1Ab (Garcia-Bates et al., 2016) at 2,000 MOI. Cells were collected after 7 d in culture for analyses of checkpoint molecules and MART1_{27–35}-specific T cells as described previously. An aliquot of cells from the in vitro cultures was used and stimulated with MART1_{27–35} peptide-pulsed T2 or Mel526 cells in the presence of 10 μ g/ml Brefeldin A (Sigma-Aldrich). After 5 h, cells were stained with CD3 (SK7), CD8 (RPA-T8), CD69 (FN50), and CD107a (H4A3; BD Biosciences) and were fixed with 1.5% paraformaldehyde (eBioscience) and subsequently exposed to 1 \times permeabilization buffer (eBioscience) before intracellular staining with IFN- γ (B27) and TNF- α (Mab11; BD Biosciences).

Multiplex gene expression analysis

Total RNA isolated from circulating lymphocytes and patient tumor biopsies (Table S4) was isolated as described previously (Vujanovic et al., 2019). The NanoString nCounter platform was used for targeted mRNA and protein expression analyses. The NanoString PanCancer Immune Profiling (770 genes) and CAR-T Characterization (780 genes) panels were used to profile gene expression in circulating lymphocytes, and the NanoString PanCancer Progression panel (770 genes plus 30 custom targets) was used for profiling of optimal cutting temperature-embedded melanoma tumor samples. The expression of 30 target proteins was analyzed using the Vantage 3D Protein Solid Tumor Panel. Raw protein count data were preprocessed by applying background thresholding and content normalization in NanoString nSolver 4.0. For gene expression analysis, the R package NanoStringNorm (Waggott et al., 2012; version 1.2.1) was used for preprocessing of the raw count data (background subtraction and normalization to housekeeping genes). Differential gene expression was analyzed using limma (version 3.38.3) with weights generated by the voom function (Law et al.,

2014; Ritchie et al., 2015). A \log_2 fold change of two and P value threshold of 0.05 were used to determine statistical significance. For outcome-based comparisons, samples were grouped into two clinical outcome categories: good (SD, NED1 [recurrence >18 mo], PR) and bad (PD, NED2 [recurrence <18 mo]). GSEA was conducted using gene sets from the MSigDB (version 6.2) in the C2 curated gene category (2005, PNAS 102, 15545–15550). Plots were generated using the R package ggplot2 (version 3.1.1) and the javaGSEA application (version 3.0; all NanoString panels are for research use only, not for use in diagnostic procedures). Patients' banked specimens were prioritized for this analysis based on tumor specimen availability and the availability of multiple blood samples that were matched to tumor time points and obtained from elutriated leukapheresis (and not processed as Ficoll PBMCs from patients who may have refused the second or third requested leukapheresis procedure) to avoid any impact of sample processing methods on the analysis.

Serum Luminex

The Human Checkpoint 14-plex kit (Procarta Plex; Thermo Fisher Scientific) was used for detection of serum checkpoint and costimulatory molecules (Butterfield et al., 2019).

Statistical analysis

Statistical comparisons between groups were performed using the Wilcoxon rank sum test, Wilcoxon signed rank test, Welch's *t* test, paired-sample *t* tests, or linear mixed effect models, as indicated in figure legends. P values are represented as *, P ≤ 0.05 , **, P ≤ 0.01 , ***, P ≤ 0.001 , and ****, P ≤ 0.0001 . P values <0.05 were considered statistically significant. The Shapiro-Wilk test was used to assess data normality, and statistical tests were performed using R (version 3.6.1). Figure graphs were generated using the R package ggplot2 (version 3.1.1). Kaplan-Meier survival curve analysis and Cox proportional-hazards modeling were performed using the R packages survival (version 3.1–8) and survminer (version 0.4.6). Standardized effect sizes were computed using the R package effsize (0.7.6), and equivalence testing was performed using the equivalence package (version 0.7.2).

Online supplemental material

Fig. S1 shows associations of MA expression levels in the DC vaccines with survival time and clinical responses. Fig. S2 shows post-trial assessment of MA-specific CD8 T cells from two HLA-A2 $^+$ patients 1.5–2 yr after DC vaccination. Fig. S3 shows immune checkpoint profiling in circulating lymphocytes and Luminex immune checkpoint profiling in serum. Fig. S4 shows differential gene expression and immune checkpoint pathway associations in circulating lymphocytes and tumor/TIL samples. Fig. S5 shows differentially expressed genes in circulating lymphocytes at d0 and d43. Table S1 lists patient demographics. Table S2 lists the statistical analyses of checkpoint and exhaustion markers. Table S3 contains statistical analyses of checkpoint and exhaustion markers CTLA-4, LAG3, PD-1, and TIM3 on CD8 T cells stimulated with DC transduced with AdVMART1 and with or without Ad5.hPD1Ab in vitro. Table S4 shows circulating lymphocyte and tumor specimens from patients used for

NanoString gene expression profiling. Table S5 shows a time-dependent covariate Cox regression analysis of soluble cytokines.

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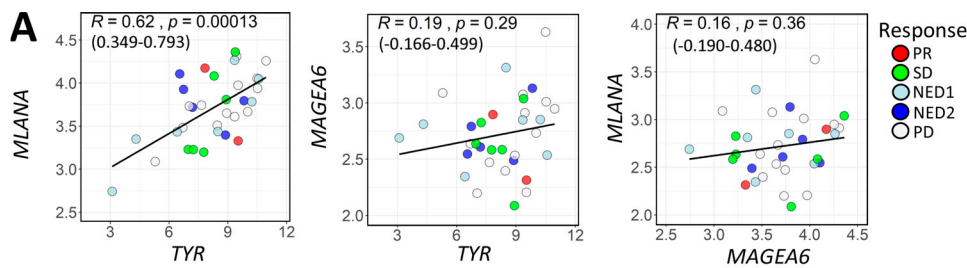
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Supplemental material



B Melanoma antigen mRNA levels associated with OS/PFS

	Factor	HR ¹	95% CI ²	p-value
OS ³ MA ⁴ mRNA expression in AdVTMM2 (n=33)	TYR	0.89	0.54-1.5	0.66
	MLANA	0.99	0.63-1.6	0.97
	MAGEA6	1.6	0.98-2.7	0.061
PFS ⁵ MA mRNA expression in AdVTMM2 (n=33)	TYR	1.1	0.75-1.8	0.54
	MLANA	0.97	0.7-1.4	0.87
	MAGEA6	1.2	0.78-1.9	0.37

¹hazard ratio, ²confidence interval, ³overall survival, ⁴tumor antigen, ⁵progression-free survival

C Associations between MA-specific PBMC IFN- γ release and OS/PFS at:

Baseline					Post-vaccine				
	Factor	HR ¹	95% CI ²	p-value		Factor	HR ¹	95% CI ²	p-value
OS ³ d0 TA ⁴ -specific PBMC responses (n=30)	Tyrosinase	1.6	0.92-2.7	0.095	OS ³ d43 TA ⁴ -specific PBMC responses (n=30)	Tyrosinase	1.4	0.83-2.4	0.21
	MART-1	1.6	0.96-2.6	0.073		MART-1	0.93	0.51-1.7	0.81
	MAGE-A6	1.4	0.76-2.5	0.28		MAGE-A6	1.6	0.99-2.5	0.057
	LacZ	0.86	0.47-1.6	0.62		LacZ	0.75	0.35-1.6	0.46
PFS ⁵ d0 TA-specific PBMC responses (n=30)	Tyrosinase	1.3	0.82-1.9	0.29	PFS ⁵ d43 TA-specific PBMC responses (n=30)	Tyrosinase	1.1	0.67-1.8	0.73
	MART-1	0.89	0.55-1.4	0.63		MART-1	1.1	0.65-1.8	0.74
	MAGE-A6	1.1	0.65-1.8	0.78		MAGE-A6	0.92	0.57-1.5	0.73
	LacZ	0.89	0.6-1.3	0.58		LacZ	1.3	0.84-2	0.24

¹hazard ratio, ²confidence interval, ³overall survival, ⁴tumor antigen, ⁵progression-free survival

Figure S1. Associations of MA expression levels in the DC vaccines with survival time and clinical responses. **(A)** Scatter plots showing correlations between TYR (Tyrosinase) and MLANA (MART-1; in the same expression cassette), TYR and MAGEA6 (MAGE-A6; separate cassettes; equivalence test for $R > -0.3$ and $R < 0.3$; $P = 0.258$) and MLANA and MAGEA6 (separate cassettes; equivalence test for $R > -0.3$ and $R < 0.3$; $P = 0.215$) based on mRNA expression measured in the DC vaccines. Pearson correlation coefficient (R), with P values and 95% confidence intervals indicated. **(B)** Univariate Cox regression analyses for DC vaccine-MA antigen expression levels and OS and PFS. **(C)** Univariate Cox regression analysis for associations between MA-specific IFN- γ release (ELISPOT counts) from total PBMCs isolated at baseline (d0) or after vaccine (d43) and OS and PFS.

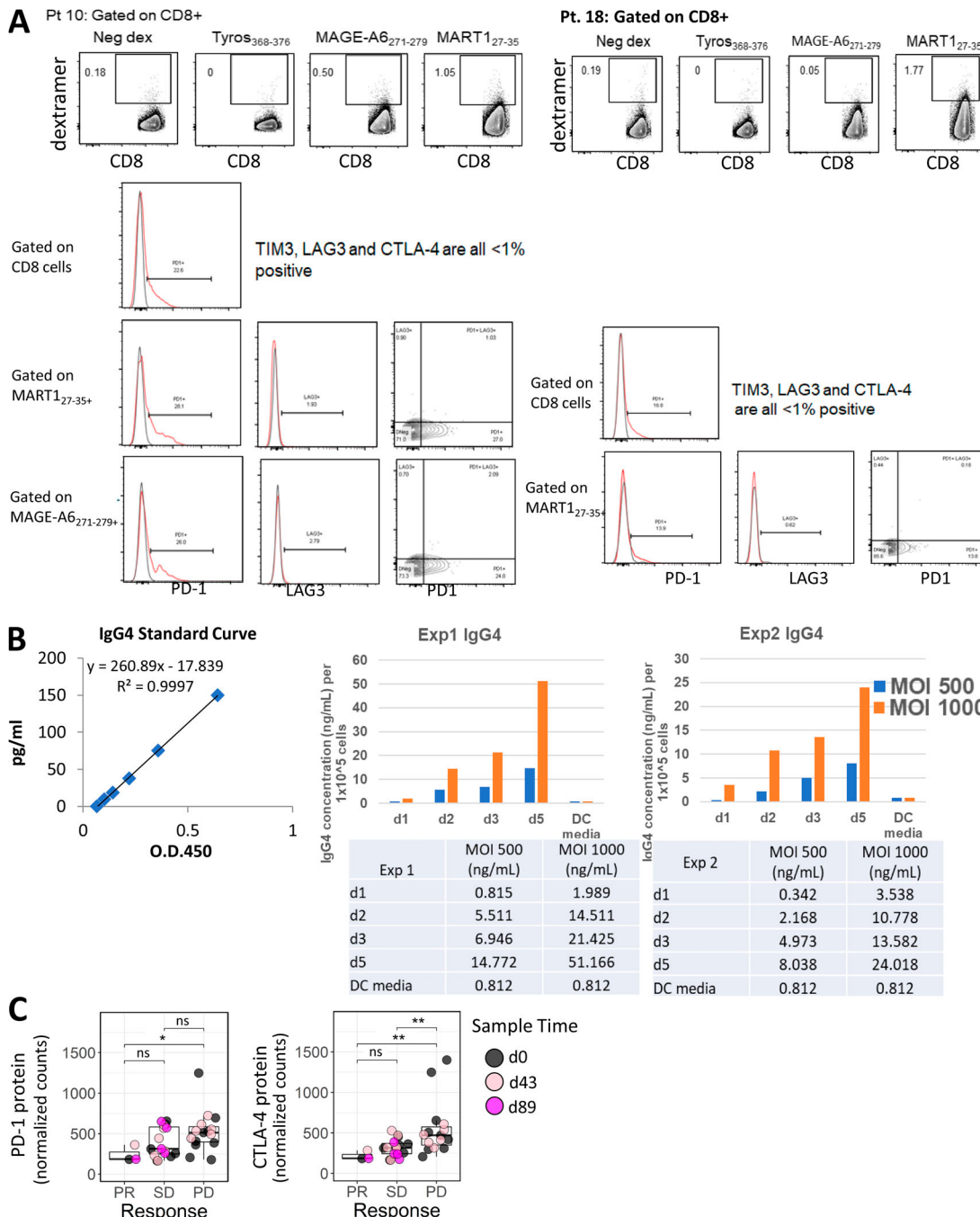


Figure S2. Post-trial assessment of MA-specific CD8 T cells from two HLA-A2⁺ patients 1.5–2 yr after DC vaccination. PBMC samples from two patients were used to determine the frequency of circulating dextramer-specific CD8 T cells as well as to examine checkpoint expression in CD8 and MA-specific CD8 T cells for patient 10 and patient 18 (A), 1.5 and 2 yr after DC vaccination, respectively. Frequencies shown for each dextramer were calculated by subtracting the negative control dextramer (Neg dex) frequency for each sample tested. Raw data histograms are shown with isotype controls for coexpression of other proteins as labeled. (B) Measurement of IgG4 in serum-free supernatants of HD DC transduced with Ad5.hPD1Ab. Two different HD monocyte-derived DC preparations were cultured for 5 d, harvested, and transduced with 500 or 1,000 MOI Ad5.hDP1Ab for 3 h before replating using serum-free AIMV media. Supernatant aliquots were taken each day as indicated after DC transduction. The concentration of human IgG4 in supernatants was quantified using a human IgG4 ELISA kit (Invitrogen) per manufacturer's instructions. (C) Graphs show distribution of PD-1 and CTLA-4 protein expression in circulating lymphocytes grouped according to clinical response. The Wilcoxon rank sum test was used for calculating P values. *, P ≤ 0.05; **, P ≤ 0.01. ns, not significant; Tyros, tyrosinase.

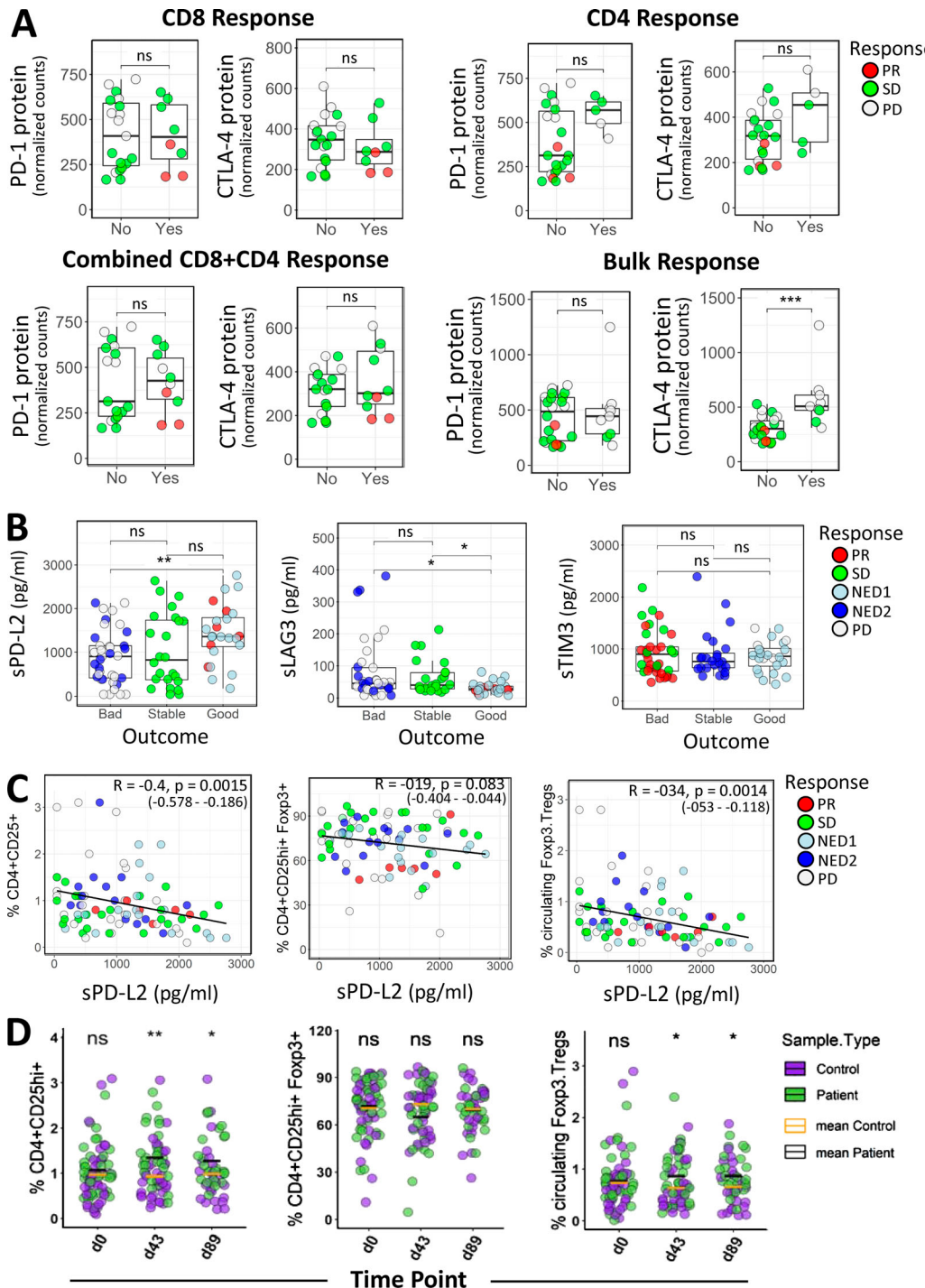


Figure S3. Immune checkpoint profiling in circulating lymphocytes and Luminex immune checkpoint profiling in serum. **(A)** Distributions of checkpoint PD-1 and CTLA-4 protein expression in relation to patient-derived MA-specific CD8, CD4, combined CD8+CD4, and Bulk (Total PBMCs) IFN- γ T cell responses, as defined in Materials and methods. **(B)** The Human Checkpoint 14-plex assay kit (Procarta Plex) was used to measure serum proteins, including checkpoint, costimulatory, and exhaustion markers in 35 patients profiled at baseline ($n = 35$), d43 ($n = 30$), and d89 ($n = 20$). Shown are distributions of soluble cytokine sPD-L2, sLAG3, and sTIM3 levels across the clinical outcome groups. For A and B, the Wilcoxon rank sum test was used for calculating P values. *, $P \leq 0.05$; **, $P \leq 0.01$; ***, $P \leq 0.001$. **(C)** Scatter plots showing correlations between FACS T reg cell profiles and sPD-L2 levels. Spearman correlation coefficient (R), P values, and 95% confidence intervals are indicated. **(D)** Percentage distributions of FACS T reg cells in melanoma patient and normal donor samples. P values denoting significance between patient and healthy samples at indicated time points were calculated using the unpaired Welch's t test. *, $P \leq 0.05$; **, $P \leq 0.01$. ns, not significant.

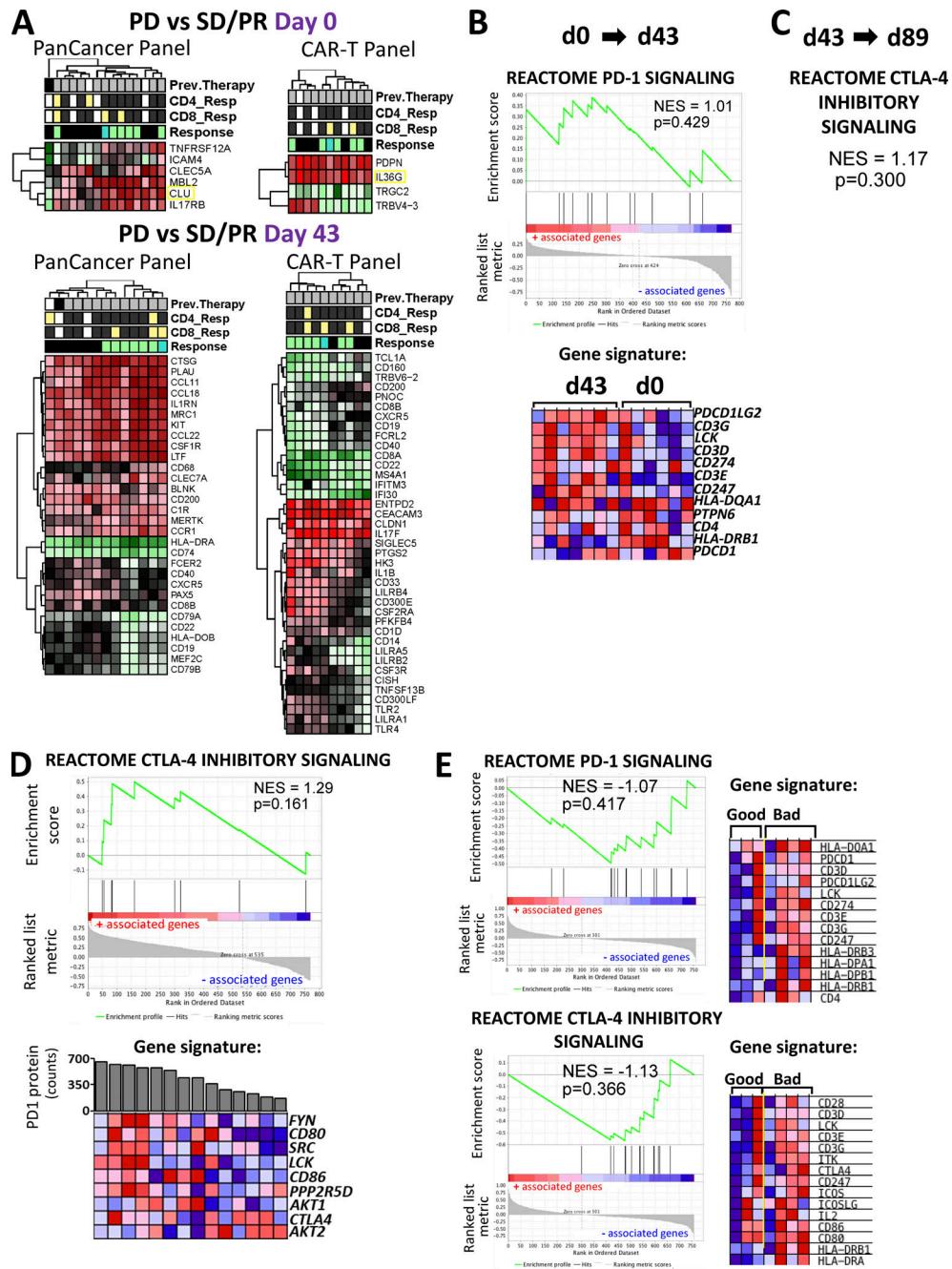


Figure S4. Differential gene expression and immune checkpoint pathway associations in circulating lymphocytes and tumor/TIL samples. (A) Heatmaps for differentially expressed genes. Shown are heatmaps for differentially expressed genes between clinical response groups that were profiled using the PanCancer Immune Profiling and CAR-T Characterization panels in baseline and d43 circulating lymphocytes. Differential gene expression analysis was conducted using the limma R package as described in the Materials and methods section. Hierarchical clustering of DEGs was conducted using the pheatmap R package. The threshold for DE genes was set to fold change ≥ 1.85 and P value ≤ 0.05 . Relevant phenotypic patient information associated with each lymphocyte sample is illustrated above the heatmaps. **(B)** PD-1 signaling pathway enrichment changes from d0 to d43 circulating lymphocytes. Shown is a GSEA plot and associated gene signature for the PD-1 signaling pathway. This figure comes from the same set of analysis and is analogous to the CTLA-4 GSEA plot in Fig. 8C. **(C)** CTLA-4 pathway enrichment changes from d43 to d89 circulating lymphocytes. Shown are statistical parameters (NES and P value) for the CTLA-4 inhibitory signaling pathway. This figure comes from the same set of analysis and is analogous to the PD-1 GSEA plot in Fig. 8D. **(D)** Results from the GSEA correlation-based enrichment analysis, which used the numeric PD-1 protein expression levels as a continuous phenotype. d43 and d89 (omitting baseline samples) circulating lymphocyte expression sets were used for this analysis. Shown is a representative GSEA plot and heatmap for genes involved in the CTLA-4 inhibitory signaling pathway, which positively correlated with PD-1 protein expression (shown as bar graph above the heatmap) in patient lymphocytes. **(E)** Immune checkpoint pathways associated with clinical outcomes in tumor biopsies. Shown are GSEA plots for PD-1 signaling and CTLA-4 inhibitory pathways enriched in tumor biopsy specimens from unfavorable (bad) clinical outcome patient groups. The associated gene signatures are presented in the form of heatmaps next to GSEA plots. Heatmaps in B-E show the clustered genes in the leading-edge subsets for each pathway category. Gene expression values are represented as colors, where the range of colors (red, pink, light blue, dark blue) shows the range of expression values (high, moderate, low, lowest). Prev., previous.

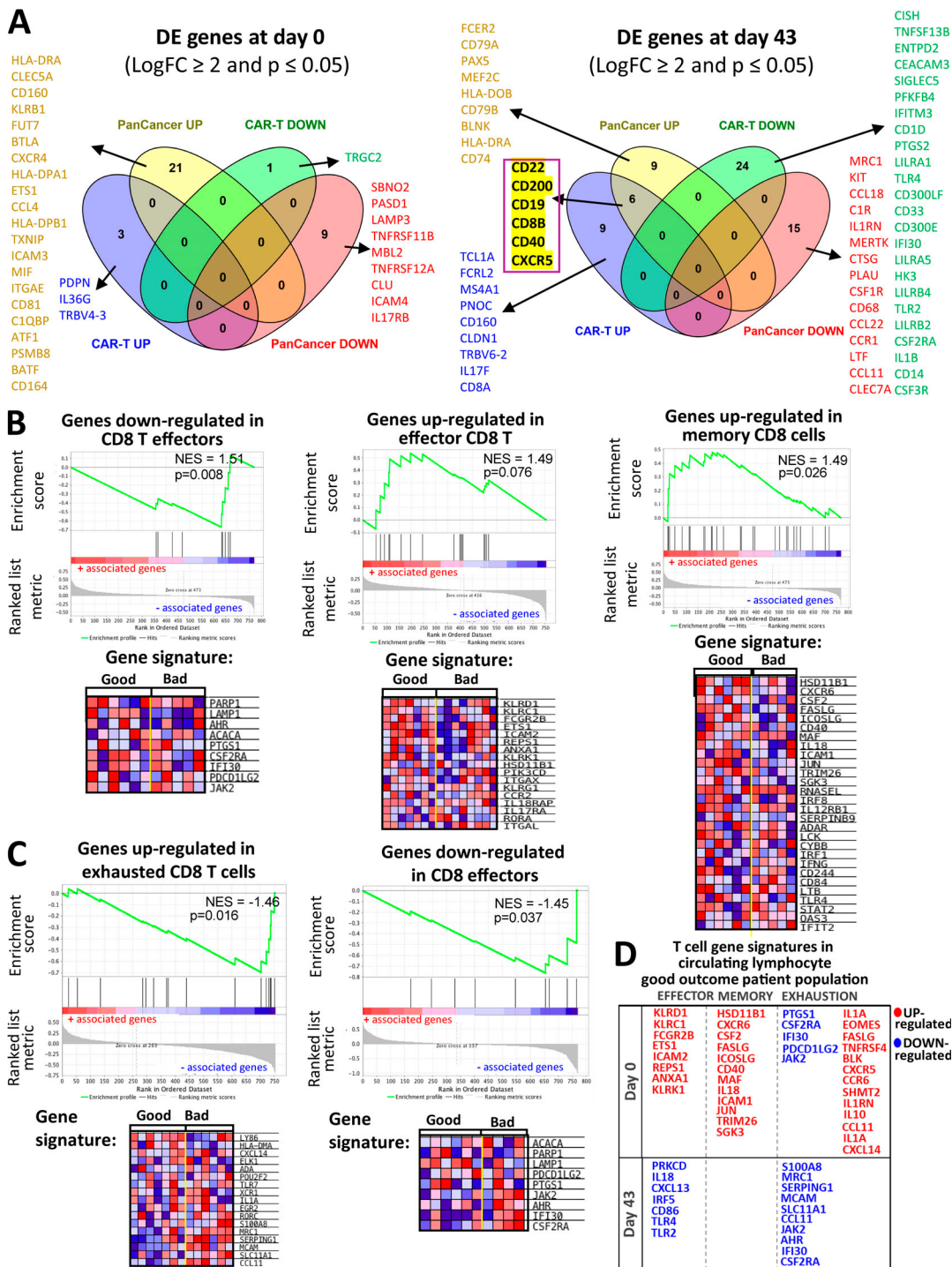


Figure S5. Comparisons for differentially expressed genes between the NanoString Panels and GSEA analysis of T cell exhaustion gene profiles.
(A) Summary of differentially expressed (DE) genes in circulating lymphocytes at d0 and d43. Venn diagrams show differentially expressed genes ($\text{LogFC} \geq 2$ and $P \leq 0.05$) in baseline (left) and d43 lymphocyte samples (right) that are represented in both the PanCancer Immune Profiling and CAR-T Characterization NanoString gene panels used in our study. PanCancer UP/DOWN and CAR-T UP/DOWN groups contain genes that are up-/down-regulated in the PanCancer Immune Profiling and CAR-T Characterization NanoString panel, respectively. Genes from each of the Venn diagram sections are listed in the figure. A set of six genes that are highlighted in yellow denote gene probes that were detected as up-regulated in both NanoString panels. **(B and C)** Immune checkpoint pathway associations with T cell exhaustion profiles in circulating lymphocytes. Representative GSEA plots for T cell exhaustion-related immune regulatory pathways correlating with good and bad clinical outcome groups in baseline (B) and d43 (C) circulating lymphocytes. T cell exhaustion signatures were obtained using the MSigDB C7 immunological gene database. Heatmaps next to the GSEA plots show clustered genes in the leading-edge subsets for each pathway category. Gene expression values are represented as colors, where the range of colors (red, pink, light blue, dark blue) shows the range of expression values (high, moderate, low, lowest). **(D)** Effector/memory T cells versus exhaustion gene signatures in the good outcome patients' circulating lymphocytes. Here, we provide a table summary for a set of differentially expressed genes at baseline (d0) and in postvaccine samples (43) that are categorized based on their association with a particular T cell phenotype (effector, memory, or exhaustion).

Tables S1–S5 are provided online as separate Word documents. Table S1 contains HLA-A2⁺ melanoma patient demographics and previous treatments. Table S2 shows a linear mixed effect model analysis of previous checkpoint therapy and MA-specific T cell responses. Table S3 contains statistical analyses of checkpoint and exhaustion markers CTLA-4, LAG3, PD-1, and TIM3 on CD8 T cells stimulated with DC transduced with AdVMART1 and with or without Ad5.hPD1Ab in vitro. Table S4 shows circulating lymphocyte and tumor specimens from patients used for NanoString gene expression profiling. Table S5 shows a time-dependent covariate Cox regression analysis of soluble cytokines.