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Anticancer vaccination is a promising approach to increase the efficacy of cytotoxic T lymphocyte–associated protein 4 (CTLA-4) and programmed death ligand 1 (PD-L1) checkpoint blockade therapies. However, the landmark FDA registration trial for anti–CTLA-4 therapy (ipilimumab) revealed a complete lack of benefit of adding vaccination with gp100 peptide formulated in incomplete Freund’s adjuvant (IFA). Here, using a mouse model of melanoma, we found that gp100 vaccination induced gp100-specific effector T cells (Teffs), which dominantly forced trafficking of anti–CTLA-4–induced, non-gp100–specific Teffs away from the tumor, reducing tumor control. The inflamed vaccination site subsequently also sequestered and destroyed anti–CTLA-4–induced Teffs with specificities for tumor antigens other than gp100, reducing the antitumor efficacy of anti–CTLA-4 therapy. Mechanistically, Teffs at the vaccination site recruited inflammatory monocytes, which in turn attracted additional Teffs in a vicious cycle mediated by IFN-γ, CXCR3, ICAM-1, and CCL2, dependent on IFA formulation. In contrast, nonpersistent vaccine formulations based on dendritic cells, viral vectors, or water-soluble peptides potently synergized with checkpoint blockade of both CTLA-4 and PD-L1 and induced complete tumor regression, including in settings of primary resistance to dual checkpoint blockade. We conclude that cancer vaccine formulation can dominantly determine synergy, or lack thereof, with CTLA-4 and PD-L1 checkpoint blockade therapy for cancer.

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Cancer vaccine formulation dictates synergy with CTLA-4 and PD-L1 checkpoint blockade therapy

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Anticancer vaccination is a promising approach to increase the efficacy of cytotoxic T lymphocyte–associated protein 4 (CTLA-4) and programmed death ligand 1 (PD-L1) checkpoint blockade therapies. However, the landmark FDA registration trial for anti–CTLA-4 therapy (ipilimumab) revealed a complete lack of benefit of adding vaccination with gp100 peptide formulated in incomplete Freund’s adjuvant (IFA). Here, using a mouse model of melanoma, we found that gp100 vaccination induced gp100-specific effector T cells (Teffs), which dominantly forced trafficking of anti–CTLA-4–induced, non-gp100–specific Teffs away from the tumor, reducing tumor control. The inflamed vaccination site subsequently also sequestered and destroyed anti–CTLA-4–induced Teffs with specificities for tumor antigens other than gp100, reducing the antitumor efficacy of anti–CTLA-4 therapy. Mechanistically, Teffs at the vaccination site recruited inflammatory monocytes, which in turn attracted additional Teffs in a vicious cycle mediated by IFN-γ, CXCR3, ICAM-1, and CCL2, dependent on IFA formulation. In contrast, nonpersistent vaccine formulations based on dendritic cells, viral vectors, or water-soluble peptides potently synergized with checkpoint blockade of both CTLA-4 and PD-L1 and induced complete tumor regression, including in settings of primary resistance to dual checkpoint blockade. We conclude that cancer vaccine formulation can dominantly determine synergy, or lack thereof, with CTLA-4 and PD-L1 checkpoint blockade therapy for cancer.

Introduction

Therapeutic blockade of the checkpoint receptors cytotoxic T lymphocyte-associated protein 4 (CTLA-4) and programmed cell death protein 1 (PD-1) on T cells can cure patients with metastatic cancer (1–3). However, many patients do not experience a clinical benefit, and significant hurdles remain in increasing the therapeutic benefit of checkpoint blockade (4–7). Recent evidence points to poor preexisting antitumor T cell immunity (8, 9) and genomic defects in IFN-γ pathway genes in tumor cells (10) as a cause for primary resistance to CTLA-4 checkpoint blockade therapy. One strategy to increase tumor-specific T cell immunity is vaccination (11), and consequently, vaccination is being explored as an avenue to increase clinical efficacy of anti–CTLA-4 and anti–PD-1 therapies (12).

However, the same landmark study (1) of 676 patients with melanoma that led to FDA approval of anti–CTLA-4 (ipilimumab, Yervoy) showed no therapeutic enhancement of anti–CTLA-4 monotherapy by concurrent vaccination with gp100 peptide in incomplete Freund’s adjuvant (IFA). Instead, addition of gp100/IFA vaccination significantly decreased the best overall response rate ($P = 0.04$) and disease control rate ($P = 0.04$) to ipilimumab, suggesting a surprising, dominant-negative effect of the vaccine on the efficacy of CTLA-4 checkpoint blockade (1). These results leave uncertain how best to combine vaccination with checkpoint blockade to increase therapeutic efficacy.

Currently there are 33 registered clinical trials in the USA alone (according to ClinicalTrials.gov) in which a cancer vaccine is combined with anti–CTLA-4 and/or anti–PD-1 checkpoint blockade (Supplemental Table 1; supplemental material available online with this article; https://doi.org/10.1172/JCI93303DS1). Of these trials, initiated by a variety of academic and industry parties in a broad range of cancers, 14 (42%) use an IFA-based vaccine, indicating how widely this formulation is applied clinically. Our data suggest that in these trials the addition of IFA-based vaccine is likely to result in no additional, and possibly reduced, clinical benefit of checkpoint blockade therapy. Conversely, our results suggest that switching to a different vaccine formulation may result in strongly increased therapeutic efficacy. These findings may have direct implications for current and future patients treated with these combination regimens.

We previously reported (13) that vaccination with gp100 peptide in IFA creates a persisting antigen depot that primes antigen-specific CD8+ T cells, which is followed by the undesirable sequestration of these cells at the vaccination site, and eventually their exhaustion and apoptosis, resulting in negligible antitumor activity. Similar results have been observed in patients with melanoma after vaccination with multiple melanoma antigen–derived pep-
tides formulated in IFA (14). Here we test the hypothesis that the inflamed, chemokine-rich vaccination site also potently sequesters the systemic pool of checkpoint blockade–induced effector T cells with vaccine-unrelated tumor antigen specificity, reducing the efficacy of CTLA-4– and PD-1–based checkpoint blockade therapy. We subsequently demonstrate that nonpersisting, virus, or water-based vaccines do not induce these deleterious effects and instead potently synergize with CTLA-4 and PD-1 checkpoint blockade therapy.

Results

Anti–CTLA-4 therapeutic activity is impaired by addition of gp100/IFA vaccination. To understand the parameters that control synergy between checkpoint blockade and anticancer vaccination, we modeled the clinical scenario of anti–CTLA-4 therapy of melanoma plus vaccination with gp100 peptide in IFA (1) by adapting the standard mouse model of anti–CTLA-4 therapy of B16 melanoma (15–17) to include vaccination with gp100 peptide in IFA. To correct for the fact that B16 melanoma progresses so rapidly that there is no time to raise gp100-specific effector T cells (Teffs) by multiple cycles of vaccination as given to the patients (1), we adoptively transferred naive, gp100-specific T cell receptor (TCR) transgenic pmel-1 (Vb13*CD90.1+CD8+) T cells. The pmel-1 T cell recognizes the gp100 peptide derived from the murine gp100 protein encoded by the pmel-17 gene, an ortholog of the melanocyte differentiation antigen gp100, which is often overexpressed in human melanomas and recognized by human CD8+ T cells (18, 19). No untreated mice or mice receiving a mock vaccine (control/IFA), or gp100 peptide in IFA (gp100/IFA) were cured, whereas 23% of the animals were cured with anti–CTLA-4 therapy (Figure 1, A and B). The addition of control/IFA vaccination to anti–CTLA-4 therapy did not affect the therapeutic impact. Interestingly, paralleling the observations in patients (1), gp100/IFA vaccination did not enhance, but significantly decreased, the therapeutic efficacy of anti–CTLA-4 therapy despite inducing a high level of gp100-specific CD8+ Teffs in peripheral blood leukocytes (PBLs) (Figure 1, C and D, P < 0.0001). Thus, gp100/IFA vaccination reduced the therapeutic efficacy of anti–CTLA-4 therapy. To both extend and replicate our finding beyond B16 melanoma, we treated E.G7.OVA, a thymoma-expressing OVA, as a model antigen. We found that the addition of OVA257–264 peptide in IFA (OVA/IFA) immunization to primed endogenous OVA-specific T cells significantly impaired anti–CTLA-4–induced therapeutic activity (7% of mice cured) compared with anti–CTLA-4 plus control/IFA (27%) or anti–CTLA-4 monotherapy (60%), OVA/IFA (0%), control/IFA (0%), or no treatment (0%) (Supplemental Figure 1, A–C, P < 0.0001). Thus, gp100/IFA-based vaccination reduced the therapeutic efficacy of anti–CTLA-4 therapy. To both extend and replicate our finding beyond B16 melanoma, we treated E.G7.OVA, a thymoma-expressing OVA, as a model antigen. We found that the addition of OVA257–264 peptide in IFA (OVA/IFA) immunization to primed endogenous OVA-specific T cells significantly impaired anti–CTLA-4–induced therapeutic activity (7% of mice cured) compared with anti–CTLA-4 plus control/IFA (27%) or anti–CTLA-4 monotherapy (60%), OVA/IFA (0%), control/IFA (0%), or no treatment (0%) (Supplemental Figure 1, A–C, P < 0.0001). This finding suggests that vaccination-induced loss of anti–CTLA-4 activity was not a peculiarity of the B16 tumor model, of pmel-1 T cells, or of gp100 peptide as a vaccine antigen.

CD8+ Teffs localize to sites of vaccination with noncognate antigen. Vaccination with gp100/IFA sets up a persistent, chronically

Figure 1. Anti–CTLA-4 therapeutic activity is impaired by the addition of gp100/IFA vaccination. (A) Experimental scheme. (B) Mice bearing 3-day-old B16 tumors received naive gp100-specific CD90.1+ pmel-1 T cells i.v., followed by s.c. vaccination with hgp100/IFA or control/IFA on day 0 and/or anti–CTLA-4 plus Gvax therapy on days 0, 3, and 6, or were left untreated. Tumor size in individual mice is shown. Numbers in parentheses represent the proportion of mice with tumor-free survival. (C) pmel-1 T cells in PBLs 7 days after vaccination. Data represent mean ± SEM, n = 5. (D) Kaplan-Meier survival curves. Data are pooled from 2 independent experiments, each with 8 to 15 mice per group.
inflamed antigen depot at the injection site, which sequesters gp100 vaccine–activated, gp100–specific Teffs, preventing them from reaching the tumor (13). We hypothesized that sequestration of Teffs at antigen-rich vaccine sites is not restricted to vaccine antigen–specific Teffs, but extends to other Teffs with specificities for antigens unrelated to those used in the vaccine. To test this hypothesis, we adoptively cotransferred luciferase–transduced OT-I Teffs and untransduced gp100-specific pmel-1 Teffs into mice vaccinated with gp100/IFA. Four days later, we observed strong accumulation of luciferase+ OT-I Teffs at the gp100/IFA vaccination site, but minimal accumulation at the tumor or saline/IFA control vaccination site (Figure 2A, P < 0.001). When we performed the converse experiment, more luciferase+ pmel-1 Teffs accumulated at the OVA257–264 (SIINFEKL) peptide in the IFA vaccination site than at the tumor or saline/IFA site (Figure 2A, P < 0.001). Thus, specific T cell reactivity to persisting vaccine antigen induces sequestration and dysfunction of Teffs that recognize cognate vaccine antigen and also of Teffs with other antigenic specificities. Given our observation in Figure 2A, we wished to evaluate therapeutic activity of adoptively transferred OT-I T cells in mice with B16.OVA after gp100/IFA or control/IFA vaccination. We found that OT-I T cells in blood showed reduced proliferation, increased expression of inhibitory cell surface markers, and loss of therapeutic activity when a distant gp100/IFA vaccination site was present (Supplemental Figure 2, A–C).

We next studied the localization of anti–CTLA-4 therapy–induced Teffs during concurrent vaccination with gp100/IFA. As reported before, some of these Teffs recognized the melanocyte
Figure 3. Exhaustion and apoptosis of sequestered anti–CTLA-4 activated CD8+ Teffs. (A) Gating strategy for naive, non–pmel-1 Teffs and pmel-1 Teffs from PBMCs (top). Mice bearing 3-day-old, s.c. B16 melanoma received CD90.1+ pmel-1 T cells, early anti–CTLA-4 therapy, and vaccination with control/IFA or hgp100/IFA. (B) Ki67 protein expression by naive CD8+ T cells, non–pmel-1 CD8+ Teffs, and pmel-1 CD8+ Teffs in PBMCs, VdLNs, and at the vaccination site 9 days after the start of therapy. pmel-1 histograms show T cells in a gp100/IFA setting. (C) PD-1/LAG-3 expression in Teffs from PBMCs, VdLNs, tumor site, and vaccination site. (D) TRP-2–specific CD8+ Teffs in VdLNs (top panel) and quantitation of IFN-γ+TNF-α+ TRP-2–specific CD8+ Teffs in PBMCs, tumor site, and vaccination site (bottom panel). (E) Fas expression on naive T cells, non–pmel-1 Teffs, and pmel-1 Teffs from VdLNs, spleen, tumor site, and at the vaccination site 9 days after the start of therapy. (F) Apoptotic cell death from VdLNs, spleen, and at the vaccination site of naive CD8+ T cells, non–pmel-1 CD8+ Teffs, and pmel-1 CD8+ Teffs 9 days after the start of therapy, as measured by flow cytometry of annexin V and 7-AAD staining. Plots are shown as mean ± SEM (n = 5, *P < 0.05 unpaired 2-tailed t test). Data shown are representative of 2 experiments.
differentiation antigen tyrosinase-related protein 2 (TRP-2) (15, 16); however, the majority of Teffs have unidentified specificities (2, 3, 20). We therefore used a recently reported methodology to specifically detect the entire polyclonal Teff pool based on CD8αCD44 hiCD11 ahi surface expression (Supplemental Figure 3A) (21, 22), as well as IFN-γ secretion in response to specific antigen stimulation (Supplemental Figure 3B), and surveyed their localization at the tumor and vaccination sites 9 days after vaccination. Anti–CTLA-4 monotherapy markedly increased anti–CTLA-4–induced non–pmel-1 CD8+ Teff number at the tumor site (Figure 2B). As expected, a fraction of these Teffs specifically recognized epitopes from the known murine melanoma antigens TRP-2 (Figure 2C), p15E (Supplemental Figure 4A), and gp100 (Supplemental Figure 4B). When gp100/IFA vaccination was added, the vast majority of anti–CTLA-4 therapy–induced non–cognate (TRP-2 and p15E) antigen–specific and cognate (gp100) antigen–specific non–pmel-1 Teffs did not reach the tumor and instead colocalized with pmel-1 Teffs at the vaccination site (Figure 2, B–D, and Supplemental Figure 4, A and B). Thus, the combination therapy resulted in lower tumor/vaccination site ratios than CTLA-4 monotherapy for total non–pmel-1 Teffs (18:1 versus 1,366:1, P = 0.001), TRP-2–specific Teffs (0.3:5.4 versus 6:1, P = 0.005), p15E–specific Teffs (4:1 versus 86:1, P = 0.003), and non–pmel-1 gp100–specific Teffs (14:1 versus 29:1, P = 0.006). Combination therapy also caused a significantly lower tumor/vaccination site ratio for gp100–specific pmel-1 Teffs (4:1 versus 36:1, P = 0.0004), resulting in reduced tumor control (Figure 2E). The gp100/IFA vaccination site also sequesters more TRP-2–specific T cells than spleen (>4-fold), vaccine-draining lymph nodes (VdLN), and tumor (>18-fold) (Supplemental Figure 4C). Thus, addition of gp100/IFA vaccination to CTLA-4 blockade diverts both vaccination-induced and anti–CTLA-4–induced tumor-specific Teffs away from the tumor and toward the vaccination site.

Exhaustion and apoptosis of sequestered anti–CTLA-4–activated CD8+ Teffs. To determine the fate of both pmel-1 and non–pmel-1 Teffs, we first analyzed Teff activation status in peripheral blood mononuclear cells (PBMCs) (Figure 3A and Supplemental Figure 3, A and B). Next, we analyzed proliferation, function, and apoptosis markers of Teffs in PBMCs, VdLN, and spleen, and at the tumor and vaccination sites. pmel-1 Teffs proliferated vigorously at all sites in mice receiving gp100/IFA, whereas non–pmel-1 Teffs proliferated less in mice vaccinated with gp100/IFA than in mice vaccinated with control/IFA (Figure 3B). Both pmel-1 Teffs and non–pmel-1 Teffs expressed multiple activation/checkpoint molecules including PD-1 and LAG3 (Figure 3C) (23). We next analyzed T-box transcription factors eomesodermin (Eomes) and T-bet, both major mediators of T cell fate and function through regulation of expression of PD-1 (24) and IFN-γ in CD8+ Teffs (25, 26). We found downregulation of T-bet and upregulation of Eomes in both pmel-1 and non–pmel-1 CD8+ Teffs at the vaccination site (Supplemental Figure 4D). We also found reduced secretion of IFN-γ and TNF-α in TRP-2–specific Teffs (Figure 3D) and p15-specific Teffs at the vaccination site (Supplemental Figure 4E). Thus, non–pmel-1 Teffs recruited to the vaccination site are exhausted at the level of surface markers, transcription factors, and cytokine production (27). In mice receiving peptide vaccination, the defect in cytokine production was maintained in the pool of Teffs that did reach the tumor site (Figure 3D). Teffs at the vaccination site also showed increased expression of Fas/CD95 death receptor (Figure 3E). Simultaneously, we observed a large population of CD11 b+Gr-1+ myeloid cells with high programmed death ligand 1 (PD-L1) and Fas ligand (FasL) expression at the gp100 vaccination site (Supplemental Figure 4F). Teffs at the vaccination site and in the VdLN showed unchanged expression of prosurvival Bcl-xL and proapoptotic Bim, but reduced expression of the prosurvival protein Bcl-2 (Supplemental Figure 4G) and a high degree of apoptotic death (Figure 3F). Together, these data indicate that pmel-1 and non–pmel-1 Teffs become sequestered at the gp100/IFA vaccination site, where they undergo proliferation, functional exhaustion, and eventual elimination through apoptotic death.

CXCR3 is required for CD8+ Teff localization to tumor. Teff migration from blood into peripheral tissues, including tumors, is regulated by interactions between chemokine receptors on Teffs and their ligands expressed in target tissues. We explored the molecular mechanism of CD8+ Teff accumulation at the tumor and vaccination sites, possibly pointing to avenues of shifting Teff accumulation from vaccination site to tumor site. To this end, we evaluated chemokine receptor and ligand expression (28–30) following therapy with anti–CTLA-4 and vaccination (Figure 4A). Among a repertoire of chemokines including 4 C-C motif receptors (CCR4, CCR5, CCR6, and CCR7) and C-X-C motif receptor 3 (CXCR3) and their canonical ligands, only CXCR3 and its ligands showed concordant expression (Figure 4B and Supplemental Figure 5A). Both pmel-1 and non–pmel-1 Teffs expressed CXCR3 in blood (Figure 4B), whereas CXCR3’s ligands, CXCL9 and CXCL10, were highly expressed at the tumor and vaccination sites (Figure 4C and Supplemental Figure 5B). Notably, the addition of gp100/IFA vaccination to anti–CTLA-4 therapy significantly reduced CXCL9 and CXCL10 expression in the tumor compared with anti–CTLA-4 monotherapy (P < 0.003). IFN-γ levels at both the tumor and vaccination sites correlated with CXCL9 and CXCL10 levels (Figure 4C), consistent with IFN-γ’s known ability to increase transcription of CXCL9 and CXCL10 in endothelial cells and myeloid cells (31, 32). Antibody blockade of CXCR3 reduced therapeutic antitumor efficacy of anti–CTLA-4–based therapy (Figure 4D), and reduced accumulation of pmel-1 and non–pmel-1 Teffs at both the tumor and vaccination sites (Figure 4E). In patients with melanoma, CXCL9 and CXCL10 expression in tumor correlated with improved overall survival, likely through the resulting infiltration of tumor-controlling CD8+ T cells (Figure 4F and Supplemental Figure 6) (33–35). Overall, Teffs induced by anti–CTLA-4 therapy and vaccination required CXCR3 to localize to both the vaccination site and the tumor site, and blockade of CXCR3 did not shift Teffs from vaccination site to tumor site.

CD8+ Teff accumulation at vaccination and tumor sites is dependent on LFA-1/ICAM-1 interaction. While chemokines are critical for T cell trafficking to tissues, T cell entry and retention also require CD8+ Teff engagement with adhesion molecules, expressed as surface glycoproteins on a variety of hematopoietic and non-hematopoietic cells (36, 37). Since leukocyte function–associated antigen 1 (LFA-1, also known as Ly49) and very late antigen 4 (VLA-4, also known as α4β1) were both expressed on pmel-1 Teffs and non–pmel-1 Teffs in blood (Figure 5B), we analyzed expression of their cognate ligands, intercellular adhesion molecule 1

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Figure 4. CXCR3 is required for CD8+ Teff localization to tumor. (A) Experimental scheme. Mice bearing 3-day-old, s.c. B16-BL6 melanomas received naive pmel-1 T cells and early anti–CTLA-4 therapy or vaccination with hgp100 in IFA. (B) Chemokine receptor expression on naive CD8+ T cells, pmel-1 CD8+ Teffs, and non–pmel-1 CD8+ Teffs in blood 9 days after vaccination. (C) Mice bearing 3-day-old, s.c. B16-BL6 melanomas received pmel-1 T cells and early anti–CTLA-4 therapy or vaccination with hgp100/IFA or control/IFA. Cytokine and chemokine concentrations in supernatant from tumor and vaccination site homogenates 9 days after vaccination. All data shown are mean ± SEM and are representative of 3 experiments (n = 5 mice per group, *P < 0.05 determined by unpaired 2-tailed t test). (D) Kaplan-Meier survival curves. (E) Absolute number of non–pmel-1 and pmel-1 T cells (mean ± SEM, n = 5) at the tumor and vaccination sites (mean ± SEM, n = 5) analyzed 9 days after the start of CXCR3 blockade. Data shown are representative of 3 experiments. (F) CXCL9 and CXCL10 mRNA expressions by RNA sequencing and overall patient survival obtained from public TCGA repositories (https://tcga-data.nci.nih.gov and http://gdac.broadinstitute.org/). Bars show mean ± SEM CD8 mRNA level as determined by paired 2-tailed t test.
(ICAM-1) and vascular cell adhesion molecule 1 (VCAM-1), on vascularization in the tumor and vaccination sites. Anti–CTLA-4 monotherapy markedly enhanced ICAM-1 and VCAM-1 expression by the tumor endothelium in an IFN-γ–dependent manner (Figure 5, A and C) (38, 39). This enhancement was markedly blunted by addition of gp100/IFA vaccination (Figure 5C), as was tumor infiltration by pmel-1 and non–pmel-1 Teffs (Supplemental Figure 7). At the vaccination sites, treatment-induced ICAM-1 and VCAM-1 expression was not dependent on IFN-γ, lacked clear association with the vasculature (Supplemental Figure 8), and instead was abundant on SSC⁺CD11b⁺Ly6G⁻Ly6C⁺F4-80⁻CCR2⁺ inflammatory monocytes (iMos) and SSC⁺CD11b⁺Ly6G⁺Ly6C⁺F4-80⁻CCR2⁻ granulocytes (Grans) (Figure 6A). To evaluate whether ICAM-1 blockade could shift Teffs from the vaccination site to the tumor site, we blocked ICAM-1 and found that therapeutic impact was significantly reduced (Figure 6B), and both pmel-1 Teffs and non–pmel-1 Teffs at the vaccination and tumor sites were reduced (Figure 6C). Interestingly, ICAM-1 blockade reduced iMos at the tumor site following anti–CTLA-4 monotherapy, and at the gp100/IFA vaccination site in vaccinated mice, whereas Grans remained unchanged (Figure 6D). iMos expressed elevated levels of ICAM-1 and VCAM-1 (Figure 6A), as well as E-selectin (CD62E) and P-selectin (CD62P) (data not shown), possibly explaining their ability to efficiently retain Teffs at the vaccination site (40, 41) (Figure 6C and Supplemental Figure 7). The Cancer Genome Atlas (TCGA) mRNA data analysis of tumor resections from patients with melanoma indicated that higher ICAM-1 expression correlated with improved survival (P < 0.01) and increased CD8⁺ T cell tumor infiltration (P < 0.0001) (Figure 6E), supporting the notion that ICAM-1 is critical to T cell–mediated tumor rejection in mice and humans. These results indicate that IFN-γ is recruited to the IFA vaccination site. iMos and other stromal cells are induced to produce chemokines, drawing in more T cells, which through IFN-γ release draw in more iMos, in a vicious cycle of myeloid- and T cell–driven inflammation.

Nonpersistent viral vaccine synergizes with therapeutic CTLA-4 and PD-L1 checkpoint blockade. Clinically used gp100/IFA vaccine formulation is poorly biodegradable and forms a persistent antigen depot in vivo (13, 14). Since we here observed that IFA-based vaccination also sequestered anti–CTLA-4–induced CD8⁺ Teffs, with specificities unrelated to the vaccine antigen, we tested whether these deleterious effects could be overcome by using a different, nonpersistent vaccine formulation. Immunization of mice with DCs pulsed with gp100 peptide significantly increased tumor regression and tumor-free survival after anti–CTLA-4 treatment (47%) compared with mice receiving anti–CTLA-4 monotherapy (13%). DCs pulsed with irrelevant chicken OVA peptide plus anti–CTLA-4 (0%), DCs pulsed with gp100 peptide (13%), DCs pulsed with OVA (0%), or no treatment (0%) (Supplemental Figure 11, A–C; P < 0.0001). Thus, anti–CTLA-4–induced tumor destruction was abrogated by vaccination with gp100 peptide in IFA, but enhanced by vaccination with DCs pulsed with gp100 peptide. Similarly, vaccination with vesicular stomatitis virus encoding gp100 (VSV.gp100) synergized with anti–CTLA-4 therapy, resulting in a significantly greater tumor-free survival rate (53%) compared with treatment with anti–CTLA-4 (22%), anti–CTLA-4 and gp100/IFA (8%), or VSV.gp100 alone (13%) (P < 0.0001, Figure 8, A and B). Since PD-L1 blockade is another major checkpoint blockade strategy for patients with cancer, we also tested vaccination in a setting of PD-L1 blockade. We observed that anti–PD-L1 therapy
Figure 5. Impact of vaccination on the ICAM-1/VCAM-1 expression on tumor vasculature. (A) Experimental scheme. (B) Expression of adhesion surface markers on naive, pmel-1, and non–pmel-1 CD8+ T cells in PBMCs 9 days after vaccination, anti–CTLA-4 therapy, or left untreated. (C) Mice bearing B16-BL6 melanomas were injected i.v. with pmel-1 T cells. Mice were immunized with hgp100 or control in IFA on day 3; and/or received anti–CTLA-4 therapy on days 3, 6, and 9; and/or were neutralized with IFN-γ on days 3, 5, 7, 9, and 11 after tumor injection; or they were left untreated. Graphs show ICAM-1 or VCAM-1 expression on CD31+ vasculature at tumor site 9 days after vaccination. Data are mean ± SEM and are representative of 2 experiments (n = 5 mice per group, *p < 0.05 determined by nonparametric Kruskal-Wallis test).
was also impaired by gp100/IFA vaccination; however, vaccination with VSV.gp100 strongly synergized with PD-L1 blockade, curing 50% of the mice (Figure 8C). Thus, vaccine formulation directly controlled the ability of vaccination to synergize with the CTLA-4 and PD-L1 checkpoint blockade. As before (Figure 4E), CXCR3 mediated the accumulation of both gp100-specific and TRP-2-specific CD8+ Teffs at the tumor site and gp100/IFA vaccination site in an in vivo trial using tumor-based vaccination (Supplemental Figure 12). Remarkably, the addition of gp100/IFA abrogated the efficacy of VSV.gp100 vaccination, sharply reducing intratumoral Teff numbers and animal survival (Supplemental Figure 12). Regardless of vaccine formulation, pmel-1 Teffs were found in multiple tissues, but only gp100/IFA, and not gp100/saline or VSV.gp100, vaccination recruited large numbers of pmel-1 Teffs to the vaccination site (Supplemental Figure 13). Over time, gp100/IFA vaccination resulted in progressively reduced peripheral blood Teff levels, proliferation, trafficking, and IFN-γ secretion (Supplemental Figure 14, A–D). Thus, T cell sequestration after gp100/IFA vaccination is a dominant effect that can abrogate effective checkpoint blockade monotherapy, or even effective combination therapy with checkpoint blockade and an otherwise effective, nonpersistent (viral) vaccine.

Synergy between anti–CTLA-4 therapy and VSV.gp100 was also impaired in genetically ICAM-1-deficient mice (Supplemental Figure 15, A and B). ICAM-1 antibody blockade decreased tumor infiltration of Teffs, and in the presence of persistent gp100/IFA vaccine, a markedly higher number of functional Teffs accumulated in the blood (Supplemental Figure 15, C and D) (45), likely due to their inability to exit the blood into target tissues. Taken together, the results indicate that ICAM-1/LFA-1 interaction is critical for CD8+ Teff accumulation in the tumor, regardless of vaccine formulation.

Nonpersistent vaccine formulations overcome primary resistance to CTLA-4 and PD-L1 checkpoint blockade therapy. Viral vaccines form one of several classes of nonpersistent anticancer vaccine formulations. It is therefore possible that the difference in therapeutic synergy between virus-based and IFA-based vaccination were not due to the latter’s establishment of a persistently inflamed vaccination site, but to unique immunogenic attributes of the viral vaccine. To directly establish the importance of antigen formulation, we compared vaccination with the persistent gp100/IFA vaccine formulation to vaccination with a nonpersistent, water-based gp100 peptide formulation in combination with anti–CTLA-4 in mice with more advanced (7-day), established melanoma (Figure 9A). Since water-based peptide vaccines do not efficiently prime T cell responses, we supplemented both the water-based and IFA-based vaccine with a combination of immunostimulatory molecules (agonistic anti-CD40 mAb, the TLR7 agonist imiquimod, and IL-2, collectively named covax) (13). After receiving the nonpersistent, water-based vaccine formulation, 21% of mice were cured compared with 6% of mice cured after vaccination with gp100/IFA (P < 0.0006, Figure 9B). We also tested synergy with PD-L1 checkpoint blockade therapy (46) and found that nonpersistent, water-based gp100 peptide vaccine activity synergizes with anti–PD-L1 blockade. In fact, 44% of mice receiving anti–PD-L1 and gp100/saline treatment were cured (>200 days) compared with mice receiving anti–CTLA-4 and gp100/saline (24%), anti–PD-L1 (0%), anti–CTLA-4 (0%), or gp100/saline monotherapy (0%) (Figure 9C). Similar synergy was seen between PD-L1 blockade and viral vaccination, whereas IFA-based vaccination significantly blunted therapeutic efficacy of PD-L1 blockade (Figure 9D).

Finally, we tested persistent versus nonpersistent vaccine formulations in combination with dual CTLA-4/PD-L1 checkpoint blockade therapy, which is currently the most potent therapy for patients with metastatic melanoma (2). In a challenging setting of 7-day established tumors where dual checkpoint blockade cured only 10% of the mice, addition of nonpersistent, viral gp100 vaccination resulted in the cure of 67% of the mice (P < 0.0001, Figure 9E). Conversely, dual checkpoint blockade with gp100/IFA vaccination did not cure any mice, and gp100/IFA vaccination completely destroyed the therapeutic efficacy of VSV.gp100 vaccination plus checkpoint blockade (Figure 9E).

Discussion

The introduction of CTLA-4 and PD-1 checkpoint blockade therapy is a milestone in cancer treatment, yet clinical benefit is limited to a subset of patients (1, 3, 47–51). Based on their mechanisms of action, CTLA-4 and PD-1 blockades are most effective in the presence of preexisting tumor-specific T cell responses (9, 52–54). However, due to poor immunogenicity of many tumors, such CD8+ T cell-mediated antitumor immunity is often weak and insufficient. Vaccination could induce expansion and diversification of the tumor-specific T cell pool, and in this regard, recent progress has been made in clinical cancer vaccine formulations, delivery, and ability to potentiate antitumor immune response (55–59). There exists a great variety of cancer vaccine formulations, each with its unique advantages (60). Clinical trials using DC vaccine (61), synthetic long peptides (62), DNA vaccine (63), RNA vaccine (64), whole-tumor cell vaccine (65, 66), and viral vaccine (67) have shown efficacy in inducing tumor-specific T cell responses, sometimes with clinical benefit. However, robust clinical evidence of increased efficacy of checkpoint blockade therapy in combination with vaccination is lacking. Here we show that peptide vaccine based on IFA, thus far the most commonly used experimental cancer vaccine formulation for combination with checkpoint blockade, failed to synergize with anti–CTLA-4 and anti–PD-L1 checkpoint blockade therapy.

Specifically, we found that tumor-specific Teffs, induced by checkpoint blockade therapy, were diverted away from the tumor by specific chemokines and adhesion molecules when a concurrent vaccine was given at a distant site, resulting in reduced antitumor activity of the checkpoint blockade therapy (Figure 10). Remarkably, these therapy-induced, tumor-specific Teffs had specificities other than the gp100 vaccine antigen, yet became sequestered at the gp100 vaccination site, where they eventually became dysfunctional and underwent apoptosis. Sequestration was mediated by engagement of chemokine receptors and adhesion molecules including CXCR3 and ICAM-1. CXCR3 and ICAM-1 controlled optimal localization of anti–CTLA-4–induced tumor-specific Teffs to both tumor and vaccination sites, and their blockade was unable to shift Teffs from the vaccination site to the tumor site. Neutralization of IFN-γ, a CD8+ T cell effector cytokine and inducer of ICAM-1 and CXCR3 ligands, actually reduced
Figure 6. CD8+ Teff accumulation at vaccination and tumor sites is dependent on LFA-1/ICAM-1 interaction. (A) Mice bearing 3-day-old, s.c. B16-BL6 melanomas received naive pmel-1 T cells, early anti–CTLA-4 therapy, and vaccination with hgp100 in IFA. Presence of ICAM-1/VCAM-1 inflammatory monocytes and granulocytes in blood and at the vaccination site 9 days after gp100/IFA vaccination. (B–D) Mice bearing 3-day-old, s.c. B16-BL6 tumors received i.v. pmel-1 T cells and vaccination with gp100/IFA and/or anti–CTLA-4 therapy, followed by anti–ICAM-1 or IgG (i.p.) on days 3, 5, 7, 9, and 11 after tumor injection. Tumor and vaccination sites were analyzed 9 days after the start of ICAM-1 blockade (B) Kaplan-Meier survival curves, 2 independent experiments each with n = 10 mice per group. *P < 0.05, log-rank test. (C) Non–pmel-1 and pmel-1 CD8+ Tefs at the tumor and vaccination sites. Data are mean ± SEM. n = 5, *P < 0.05 as determined by unpaired 2-tailed t test. (D) iMos and Grans at the tumor and vaccination sites. Plots are mean ± SEM, n = 5, *P < 0.05 as determined by unpaired 2-tailed t test. Data shown (B–D) are representative of 2 experiments. (E) ICAM-1 mRNA expression by RNA sequencing and overall patient survival obtained from public TCGA repositories (https://tcga-data.nci.nih.gov and http://gdac.broadinstitute.org/).
CD8+ Teff accumulation at the tumor, shifting Teffs to the vaccination site. Our findings serve to illustrate that the expression of ICAM-1 and CXCR3 ligand molecules is crucial to facilitate Teff adhesion in both the tumor and other inflamed tissues, including vaccination sites. IFN-γ-induced expression of ICAM-1 and CXCR3 ligands is also markedly increased and required for therapeutic efficacy after anti-CTLA-4 monotherapy. Consequently, neutralization of these molecules is not a viable strategy to redistribute Teffs from the vaccination site to the tumor.

The presence of immune cells in human melanomas correlated with the expression of defined chemokine subsets (29, 68). Analysis of 225 primary melanomas (35) revealed overexpression of CXCR3 ligand molecules CXCL9 and CXCL10 around a high density of tumor high endothelial venules. Indeed, CXCR3 expression on tumor-infiltrating, activated T cells is considered a good prognostic marker and is associated with improved survival in patients with melanoma (34, 69) and patients with advanced serous ovarian cancer (70), suggesting that CXCL9 and CXCL10 dominantly control the localization of such activated, potentially tumoricidal T cells. TCGA data on stage I-IV human melanoma samples confirm that mRNA levels of CXCR3, ICAM, and IFN-γ in tumors also strongly correlate with enhanced patient survival, and with the presence of CD8+ T cells. In contrast, analysis of cellular infiltrates in skin resections from vaccine injection sites in patients with melanoma following vaccination with multiple peptides in IFA showed accumulation of CXCR3+ CD8+ T cells, with minimal IFN-γ production (14, 68, 71).

Together, our data are consistent with a scenario where circulating Teffs infiltrate tumor tissue or the vaccination site and encounter cognate antigen, causing the local release of IFN-γ, which increases expression of chemokines such as CXCL9 and CXCL10 and adhesion molecules such as ICAM-1 by host cells. Increased expression of CXCL9, CXCL10, and ICAM-1 in turn attracts additional CXCR3+ Teffs, supporting a vicious cycle of T cell infiltration, cytokine secretion, and local inflammation. When the vaccine is persistently present, as in the case of gp100 peptide emulsified in nonbiodegradable IFA, this process results in chronic sequestration of Teffs at the vaccination site (13). Importantly, non-gp100-specific Teffs induced by CTLA-4 checkpoint blockade also become sequestered through CXCR3 and ICAM-1 engagement.

At the IFA vaccination site, these detrimental effects on Teffs appear to be further compounded through a distinct pattern of cooperative, chemokine-linked, feed-forward trafficking involving Teffs, iMos, and Grans. CXCR3-dependent Teff accumulation at the vaccination site drove expression of CCL2 and influx of inflammatory myeloid cells expressing ICAM-1 and VCAM-1, driving both additional T cell accumulation and inflammation in a feed-forward loop (31, 72, 73). It is possible that this same feed-forward cooperation involving (a) the link between CXCL9/10 and CCL2, and (b) LFA-1/ICAM-1-mediated cell-cell adhesion could explain why Teffs favor the vaccination site over the tumor site, which has a paucity of myeloid cells when a distant inflamed vaccination site is present. A recent study of experimental autoimmune encephalomyelitis (EAE) demonstrated that autoreactive T cells withstand the shear force of the cerebrospinal fluid and maintain their attachment to leptomeninges through LFA-1- and VLA-4-mediated cell-cell adhesion to leptomeningeal macrophages; this interaction was further potentiated by CXCR3 signaling and antigenic stimulation (74). Thus, under inflammatory conditions there appears to be an interdependent recruitment of Teffs and iMos to inflamed tissues (75). Gran recruitment to the vaccination site, on the other hand, appears to be independent of Teff recruitment, as evidenced by minimal impact of CXCR3 blockade-mediated reduction in T cell accumulation on their recruitment. CCL2 depletion reduced the number of Teffs and iMos trapped at the vaccination site, which interestingly also modestly shifted Teffs from vaccination site to tumor site. Several studies have demonstrated chemokine-mediated, cooperative trafficking of T cells and innate immune cells (31, 72, 73). In patients with melanoma, CXCR3+ T cells have also been reported to preferentially accumulate at sites of IFA-based vaccination (14). Thus, the dynamics of chemokine-linked, feed-forward cooperation among inflammatory immune cell subsets reported in the current study provide mechanistic insight into the observed increased accumulation of Teffs and inflammatory myeloid cells at cutaneous vaccination sites in mice and possibly in patients with cancer (68, 71).

Cooperation between CD8+ Teffs and iMos is increasingly regarded as a key process in several inflammation-driven human disease models (43, 75, 76). In patients with chronic obstructive pulmonary disease (COPD), inflammatory macrophages and CD8+ T cells showed enhanced migration into CXCR3 and CCR5 chemokines, leading to inflammation of the lung parenchyma and the airways (76). In patients with diabetes, CD8+ Teffs infiltrating adipose tissue exacerbated the inflammatory reaction by inducing accumulation and activation of macrophages (75). Our data suggest that a similar process also promotes detrimental T cell sequestration at the inflamed vaccination site after IFA-based vaccination. Mechanistically, we find that several critical elements of the molecular machinery of chemokines and integrins that control T cell trafficking are shared between the tumor and vaccination sites, and hence these are not suitable targets for shifting Teffs from vaccination site to tumor site. Instead, preventing the formation of a persistent, chronically inflamed vaccination site by altering vaccine formulation offers a direct path toward the design and use of vaccines that synergize with checkpoint blockade therapy.

IFA-based, long-term persisting vaccines continue to be widely used in clinical trials of cancer vaccines. However, less persistent vaccine formulations are gaining in popularity, including water-based, cellular, viral, and nucleic acid-based formulations. We previously demonstrated the utility of a water-based peptide vaccine, and here extend those findings with DC and virus-based vaccines, demonstrating marked therapeutic activity and profound synergy with checkpoint blockade. Thus, we envision a shift from long-term, persisting antigen depot vaccines to quickly resolving, short-term depot vaccines for the induction of therapeutic antitumor T cell responses.

Overall, our results indicate that IFA-based vaccination induces an inflamed vaccination site that recruits, functionally impairs, and eventually destroys tumor-specific Teffs induced by anti-CTLA-4 checkpoint blockade therapy through a mechanism dependent on inflammatory monocytes, CCL2, IFN-γ, CXCR3, and ICAM-1, causing reduced tumor control. Nonpersistent vaccine formulations can reverse these undesir-
Figure 7. CCL2 and CXCR3 mediate a feed-forward loop of CD8+ Teffs and inflammatory monocyte accumulation at the vaccination site. (A and B) Mice bearing 3-day-old, s.c. B16-BL6 melanomas were injected with naive pmel-1 T cells, vaccinated with hgp100/IFA, and received early anti–CTLA-4 therapy. (A) Distribution of CD11b+ leukocyte subsets at tumor and vaccination sites. (B) CCR2 expression on leukocyte subsets as in A and CD8+ non–pmel-1 and pmel-1 Teffs in blood observed 9 days after vaccination. (C) Mice bearing 3-day-old, s.c. B16-BL6 melanomas received pmel-1 T cells and early anti–CTLA-4 therapy or vaccination with hgp100/IFA or control/IFA. CCL2 and IFN-γ chemokine concentrations (mean ± SEM) in supernatant from tumor and vaccination site homogenates 9 days after vaccination. Data shown are representative of 3 experiments, each with n = 5 mice per group. *P < 0.05 as determined by unpaired 2-tailed t test. (D and E) Mice treated the same as in C underwent CCL2 depletion or received IgG on days 3, 5, 7, 9, and 11 after tumor injection. (D) Absolute number of iMos and Grans at the tumor and vaccination sites. (E) Absolute number of CD8+ Teffs at the tumor and vaccination sites. (F and G) Mice treated as in C received anti–CXCR3 mAb therapy or IgG on days 3, 5, 7, 9, and 11 after tumor injection. Plots shown are absolute number of (F) iMos and (G) Grans at the tumor and vaccination sites. Data shown are mean ± SEM, representative of 3 experiments, each with n = 5 mice per group. *P < 0.05 as determined by unpaired 2-tailed t test.
able effects and synergize with anti-CTLA-4 and anti-PD-L1 checkpoint blockade, overcoming primary resistance to checkpoint blockade therapy and greatly enhancing complete cure rates. These results provide a direct explanation for the puzzling lack of synergy between anti-CTLA-4 therapy and gp100/IFA vaccination in patients with melanoma. They also highlight the importance of vaccine formulation in future clinical trials of combination therapy with checkpoint blockade and vaccination and point the way to increase the therapeutic efficacy of checkpoint blockade for patients with cancer.

Methods

Mice and tumor cells. pmel-1 TCR transgenic mice on a C57BL/6 background (The Jackson Laboratory) were crossed with CD90.1 congenic mice to yield pmel-1+/− × CD90.1+/− mice (herein referred to as pmel-1 mice). C57BL/6-Tyr-2J/J albino, OT-I TCR transgenic, and ICAM-1–knockout mice were purchased from The Jackson Laboratory. B16-BL6 mice). C57BL/6J-Tyr-2J/J albino, OT-I TCR transgenic, and ICAM-1–knockout mice were purchased from The Jackson Laboratory. B16-BL6 melanoma cells and granulocyte-macrophage colony-stimulating factor (GM-CSF) producing B16-BL6 were cultured in DMEM supplemented with 1 μg/ml penicillin, 1 mg/ml streptomycin, 50 mg/ml gentamicin, 2 mM l-glutamine, and 8% FCS (Invitrogen). E.G7.OVA cells were main-

Figure 8. Nonpersistent viral vaccine synergizes with therapeutic CTLA-4 and PD-L1 checkpoint blockades. (A) Experimental scheme. Kaplan-Meier survival curves in mice with 3-day-old, s.c. B16 melanomas after receiving naive pmel-1 T cells (i.v.). (B) Early anti-CTLA-4 therapy and vaccination with hgp100/IFA (s.c.), VSV.gp100 (i.v.), VSV.OVA, or combination. (C) Early anti-PD-L1 (i.p.) plus Gvax therapy and vaccination with hgp100/IFA (s.c.), VSV.gp100 (i.v.), or combination. Data were pooled from 2 independent experiments, each with n = 10 mice per group. *P < 0.05, log-rank test.

DC generation and transfer. Bone marrow cells isolated from femurs and tibiae of C57BL/6 mice were RBC lysed and cultured in RPMI containing 10% FCS, 1 mM sodium pyruvate, HEPES, penicillin, streptomycin, and β-mercaptoethanol (EMD Millipore), supplemented with 0.1 μg/ml GM-CSF (PeproTech). On day 3, fresh RPMI containing 0.1 μg/ml GM-CSF was added. On day 7, nonadherent DCs were cultured in medium containing 0.1 μg/ml LPS for 24 hours. For transfer, DCs were pulsed for 1 hour with 10 μg/ml hgp10025–33, or chicken OVA257–264 peptides for 1 hour at 37°C. Cells were then harvested from plates and washed 2 times in PBS, and 1 × 10^6 cells per group were administered through tail vein injection.

Peptide vaccination. The synthetic, high-affinity H-2D^d-restricted heteroclitic mouse gp100227–235 peptide (hgp100, KVPRNQDWL), H-2K^d-restricted peptides chicken OVA257–264 (SIINFEKL), TRP-2240–248 (SVYDFFVWL), and p15E (KSPWFTTL) were purchased from PepTides International at a purity greater than 95%. H-2K^d-restricted TRP-2 pentamer (SVYDFFVWL) was purchased from ProImmune. Mice received 1,000 naive pmel-1 or OT-I T cells i.v. and were vaccinated with 2 separate s.c. injections at the base of the tail or in each flank with 100 μl of either saline or saline and IFA emulsion (1:1, vol/vol), each containing 100 μg hgp100227–235 or 50 μg OVA257–264 peptide. In the early treatment approach, mice were vaccinated on day 3 after tumor injection at the posterior flank with 2 injections of 100 μl IFA emulsions containing 100 μg hgp100227–235 peptide, or without peptide (control), or mice were infected i.v. with 100 μl of 1 × 10^8 PFU/ml VSV. gp100 or VSV.OVA. In the late-treatment approach, mice were vaccinated on day 7 after tumor injection at the posterior flank with 2 injec-

### Results

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erin injection. Regions of interest from the displayed images were designated and quantified as total photon counts or photons/second (represented by color bars) using Living Image software (Xenogen).

**Tissue preparation.** Mice were euthanized by CO2 inhalation. PBMCs were collected by tail bleed or cardiac puncture. Spleens, lymph nodes, and skin sections from vaccination sites and tumors were harvested and stored in cold PBS (Life Technologies). To determine the absolute number of a given cell subset per tumor or vaccination site, we adjusted cell counts quantitated by flow cytometry to absolute weight of tumor and skin from vaccine injection sites. Single-cell suspensions were prepared in PBS with 10% FCS and 2 mM EDTA (Sigma-Aldrich) by mashing tissue against the surface of a 40-μm cell strainer using the plunger of a 3-ml syringe (Becton Dickinson). RBC were removed using a hypotonic lysis buffer (StemCell Technologies). Skin sections were minced using a surgical blade (Becton Dickinson) and digested with collagenase (Sigma-Aldrich) for 3 hours at 37°C and then passed through an 18-gauge syringe (Becton Dickinson) and filtered using a 40-μm cell strainer (BD Falcon). Mice were tail bled at the indicated time points.

**Preparation of tissue homogenates.** On day 12 after tumor injection, tumor and skin sections from the vaccination site (6-mm diameter) were surgically removed from euthanized mice. Tissue weight measurements were performed. Tissues were homogenized in 1 ml cold PBS using a glass homogenizer. The homogenates were transferred into 100 μl IFA emulsions containing 100 μg hgp10025–33 peptide, or mice received covax (13) consisting of 100 μl, 1× s.c., 500 μg/ml of CD40-specific mAb (clone FGK4.5, Bioxcell), and 25 mg, 1× topical imiquimod cream 5% (Aldara, Fougera), and a total 100 μl, 5× i.p., 1 × 10^6 IU/ml rhIL-2 (TECIN, Hoffman La Roche Inc.) on days 0, 1, and 2.

**T cell imaging studies.** C57BL/6J-Tyr-2J/J albino mice were injected with 3 × 10^5 B16.white cells on the shaved back. This combination of white mice and nonpigmented tumor reduces photon absorption by the skin and tumor pigment during T cell imaging (77). Seven days later, mice were vaccinated with 100 μl saline in IFA (control) or with IFA emulsions containing 100 μg hgp10025–33 Peptide (s.c.); a 50:50 mixture of 6-day-cultured 1 × 10^6 pmel-1 T cells and 1 × 10^6 eGFP-sorted, v-effLuc–GFP-transduced OT-1 T cells (i.v.); and they received 3 days of IL-2 treatment. We performed the reverse experiment with saline in IFA (control) or IFA emulsions containing 50 μg OVA257–264 peptide (s.c.); a 50:50 mixture of 6-day-cultured 1 × 10^6 OT-1 T cells and 1 × 10^6 eGFP-sorted, v-effLuc–GFP-transduced pmel-1 T cells (i.v.); and IL-2 treatment for 3 days.

Bioluminescence imaging was performed with a cooled charge-coupled device (CCD) camera mounted in a light-tight specimen box (IVIS 200, Xenogen). For in vivo imaging, mice were given 150 mg/kg body weight i.p. of D-luciferin substrate (Caliper Life Sciences), anesthetized with 1%–3% isoflurane, and imaged 8 minutes after D-luciferin injection. Regions of interest from the displayed images were designated and quantified as total photon counts or photons/second (represented by color bars) using Living Image software (Xenogen).
to 1.5-ml Eppendorf tubes and centrifuged at 13,000 g for 10 minutes at 4°C, and the supernatant was stored at −80°C until analyzed by LumineX (Millipore) according to the manufacturer’s protocol.

**Histology.** Tumor and skin sections dissected from a similarly treated parallel group of mice as described in the previous paragraph were snap frozen in OCT. Tissues were cut into 6-μm sections then fixed with acetone/ethanol and blocked with PBS/5% BSA, anti-CD16/32, 3% H₂O₂, 0.1% NaN₃, and Avidin/Biotin Blocking Kit (Vector Laboratories). Sections were stained with biotinylated or fluorescence-conjugated antibodies (5–20 μg/ml) to CD31 (clone 390), ICAM-1 (clone YN1/1.7.4), VCAM-1 (clone 429), CD8α (clone 53-6.7), Thy1.1 (clone H1S51), and F4/80 (clone BM8) antibodies followed by peroxidase or fluorescent Streptavidin conjugates (Jackson Immunoresearch Laboratories Inc.). VCAM signals were amplified with Tyramide Signal Amplification Plus kits (PerkinElmer). Slides were mounted in SlowFade Gold (Invitrogen) and images collected on an AxioImager with Apotome (Zeiss). ImageJ (NIH) was used to analyze percent marker-positive vasculature.

**TCGA analysis.** TCGA mRNA expression by RNA sequencing and overall patient survival obtained from public TCGA repositories (TCGA Data Portal, https://tcga-data.nci.nih.gov and Broad Institute, http://gdac.broadinstitute.org/) were used to construct Kaplan-Meier curves for patients of all stages or with stage III cutaneous melanoma. When analyzing the additive effects of 2 genes on survival, we created 4 expression groups based on quartiles of median values.

**Antibody staining and FACS analysis.** Intracellular IFN-γ staining was performed using the Cytotof/Cytoperm kit (BD Biosciences) according to the manufacturer’s recommendation after 4 hours of stimulation with 1 μM mouse hgp100, or TRP-2 or p15E peptides and using a 1:800 dilution of mAb to IFN-γ (clone XMG1.2, BD Pharmingen). Annexin V binding assay was performed on unfixed cells with an annexin V–fluorescein isothiocyanate (FITC) staining kit according to the manufacturer’s instruction (BD Biosciences). All antibodies were used at 1:200 dilution unless otherwise indicated. Antibodies used for staining included antibodies against CD8 (clone 53-6.7), CD90.1 (clone HIS51), Vα2 (clone B20.1), Vβ5 (clone MR9-4), CD44 (clone IM7), CD11a (clone M17/4), TNF-α (clone MP6-XT22), CD152 (clone UC10-4B9), CD95 Fas (clone 15A7), CD11c (clone HL3), CD11b (clone M1/70), Ly-6G (clone 1A8), CD19 (clone MB19-1), CCR7 (clone 4B12), CCR6 (clone sirx6), CCR5 (clone T21/8), K-67 (clone SOAI5), Eomes (clone Dan11mag), T-bet (clone 4B10), CD54 (clone YN1/1.7.4), CD106 (clone 429), CD178 FasL (clone KAY10), and Ly-6C (clone AL-21) (all from Affymetrix/eBioscience). Antibodies against PD-L1 (10F.9G2), LAG-3 (clone C9B7W), PD-1 (clone RMPL30), CXC3 (clone CXCR3-173), CD62P (clone RMP-1), CCR6 (clone 29-2L17), CCR4 (clone 2G12), and CCR2 (clone SA203G11) were all from BioLegend. Antibodies against CD178 FasL (clone 10F.9G2), CD106 (clone 429), CD178 FasL (clone KAY10), and Ly-6C (clone AL-21) were all from BD Biosciences.

**Statistics.** All results are expressed as mean ± SEM. Mouse and sample group sizes were n = 5, unless otherwise indicated. Data were analyzed using paired or unpaired 2-tailed t tests or nonparametric Kruskal-Wallis test where applicable, and differences were considered significant at P < 0.05. All experiments were performed at least twice, with comparable results. Mice were euthanized when tumor size
reached ≥200 mm². Survival curves were plotted using Kaplan-Meier estimates and compared by log-rank analysis.

**Study approval.** Animal experiments performed in this study were approved by the Institutional Animal Care and Use Committee of the University of Texas MD Anderson Cancer Center.

**Author contributions**

YH, AW, TF, QH, MCN, FH, ZX, CV, HK, M Singh, M Sharma, FF, DM, and ZD performed experiments. SMA, KSS, TF, and PS provided study materials. AW and VHE performed immunofluorescence analysis and interpreted the results. JR performed TCGA data analysis. YH and WWO designed experiments and wrote the manuscript.

**Acknowledgments**

The work was supported by the University of Texas MD Anderson Cancer Center Specialized Program of Research Excellence in Melanoma Career Development Award (PS0 CA093459) to YH, and a Cancer Prevention and Research Institute of Texas (CPRIT) grant (RP140522) to WWO.

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