Monocyte-Derived Dendritic Cells Are Essential for CD8+ T Cell Activation and Antitumor Responses After Local Immunotherapy

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Tumors harbor several populations of dendritic cells (DCs) with the ability to prime tumor-specific T cells. However, these T cells mostly fail to differentiate into armed effectors and are unable to control tumor growth. We have previously shown that treatment with immunostimulatory agents at the tumor site can activate antitumor immune responses and is associated with the appearance of a population of monocyte-derived DCs (moDCs) in the tumor and tumor-draining lymph node (dLN). Here, we use depletion of DCs or monocytes and monocyte transfer to show that these moDCs are critical to the activation of antitumor immune responses. Treatment with the immunostimulatory agents monosodium urate crystals and Mycobacterium smegmatis induced the accumulation of monocytes in the dLN, their upregulation of CD11c and MHCII, and expression of iNOS, TNFα, and IL12p40. Blocking monocyte entry into the lymph node and tumor through neutralization of the chemokine CCL2 or inhibition of colony-stimulating factor-1 receptor signaling prevented the generation of moDCs, the infiltration of tumor-specific T cells into the tumor, and antitumor responses. In a reciprocal fashion, monocytes transferred into mice depleted of CD11c+ cells were sufficient to rescue CD8+ T cell priming in lymph node and delay tumor growth. Thus, monocytes exposed to the appropriate conditions become powerful activators of tumor-specific CD8+ T cells and antitumor immunity.

Keywords: dendritic cells, T cells, monocyte-derived dendritic cells, CSF1R, tumor immunotherapy, mouse models

INTRODUCTION

Dendritic cells (DCs) are critical for the induction of adaptive immune responses. DCs in tumors and tumor-draining lymph nodes (dLNs) are often loaded with tumor material and are able to induce proliferation of tumor-specific T cells in the dLN (1–3). Although presentation of tumor antigen by specific DC subsets is necessary for tumor regression in patients and mouse models (4, 5), it...
is mostly insufficient: proliferating T cells remain in the LN, fail to differentiate into fully armed effector cells, and are unable to infiltrate the tumor and mediate tumor rejection (3).

Several studies have shown that immunostimulating treatments that activate DCs also lead to induction of powerful CD8+ T effector cells that eliminate tumor cells and delay tumor growth (6–10). Unexpectedly, by carrying out a detailed study of the effects of these treatments on DC phenotype, we found that successful therapies do not simply activate existing DC subsets, but they also elicit the differentiation of monocytes into monocytic-derived DCs (moDCs) in the dLNs (9, 11, 12). Whether moDCs are simply passengers in the induced inflammatory response or powerful antigen-presenting cells that are especially suited to stimulating potent antitumor immunity remains unclear.

Most of our knowledge of moDCs comes from infection models. Under inflammatory conditions, moDCs become a substantial population that complements the range of steady state DCs (13–15). MoDCs upregulate CD11c and MHCII but generally retain expression of monocyte markers such as Ly6C, Ly6B, CD64, and FcRRI (15–17). They can mediate effector functions via their production of the Th1 cytokines TNF-α, CD8, and Fcε (15–17). Importantly, moDCs can also prime Th1 immunity and cytotoxic T cell responses (14, 15, 19), recognized components of successful antitumor responses (4, 20, 21).

In a tumor context, the role of monocytes as precursors of immunostimulatory moDCs is rarely considered. Instead, Ly6C+ CCR2+ monocytes are known to give rise to the majority of tumor-associated macrophages (22–24) and contribute to the pool of monocytic myeloid-derived suppressor cells (MDSCs) that promote tumor growth and suppress antitumor immunity (25, 26). Nonetheless, experiments in mouse models show that monocytes can also differentiate into inflammatory DCs that produce reactive oxygen species (ROS) and directly mediate antitumor responses independently of effector T cells (27). Similarly, in human skin lesions treated with toll-like receptor 7/8 agonists, inflammatory DCs acquire expression of cytotoxic mediators in vivo and become directly tumoricidal in vitro (28).

In addition to acquiring direct tumoricidal function, moDCs can also serve as antigen-presenting cells in the tumor context. In a recent study, anthracyclin chemotherapy induced Ly6C+ CD11c+ cells at the tumor site by an ATP- and CCR2/CCL2-dependent mechanism (29, 30). These cells, but not PDCA+ pDCs or BATF3-dependent DCs, could activate T cells in situ and were necessary for antitumor activity (29). Interestingly, lymph nodes or tertiary lymphoid tissues were not necessary for this response, suggesting that it might rely on existing memory T cells rather than de novo priming of naive T cells. It remains to be determined whether the key role of moDCs is specific to chemotherapy, or whether it may extend to antitumor immune responses induced by other treatments. In this respect, it is noteworthy that work from our group indicates that moDCs can also be elicited by peritumoral treatment with the toll-like receptor 3 ligand polyI:C, or the immunostimulatory agents monosodium urate (MSU) crystals and *Mycobacterium smegmatis* (Msmeg). Importantly, we find that the presence of moDCs correlates with CD8+ T cell activation and treatment success in several tumor models (9, 12). Whereas moDCs are frequently observed after exposure to infectious stimuli, the precise conditions leading to their generation have not been determined. In our model, polyI:C and MSU + Msmeg both induced elevated levels of IFNγ and IL-12p70 in serum (9). These cytokines were reported to play a key role in moDC generation and function (18). In addition, we observed that MSU + Msmeg induced the release of IL-1β, which was required for the antitumor response (11).

In this paper, we sought to determine the relevance of moDCs in antitumor immunity. We used a model of murine melanoma and local treatment with MSU + Msmeg to show that moDCs are critical for treatment success. MSU + Msmeg induced recruitment of monocytes from blood and their differentiation into inflammatory moDCs in the dLN. Treatment with a colony-stimulating factor-1 receptor (CSF1R) inhibitor similar to the ones currently in clinical trial to block MDSCs also blocked monocyte and moDC accumulation in the dLN, as well as tumor-specific T cell proliferation in dLN and antitumor activity. Finally, adoptively transferred monocytes were able to differentiate into CD11c+ moDCs in vivo and were sufficient to restore MSU + Msmeg antitumor responses in CD11c-depleted animals. Taken together, these results indicate that moDCs are critical for the success of MSU + Msmeg immunotherapy and suggest a common mechanism by which immunotherapy and chemotherapy may be able to transform tumors into sites of immune activation.

**MATERIALS AND METHODS**

**Mice**

All mice were bred at the Malaghan Institute of Medical Research Biomedical Research Unit. C57BL/6 (CD45.2+), B6.SJL-Ptprc+ (CD45.1+), and CD11c-DTR mice were originally from Jackson Laboratories, USA; OT1 mice expressing a transgenic TCR specific for K+ ovalbumin (OVA)257–264 were from Melbourne University, Australia. CD11c-DTR bone marrow (BM) chimeras were generated as described (9) by irradiating (2 × 550 rad) C57BL/6 hosts followed by i.v. transfer of 106 CD11c-DTR BM cells. Chimeras were rested for at least 8 weeks before being used in experiment. All experimental procedures were approved by the Victoria University of Wellington Animal Ethics Committee.

**Tumor Cell Lines and Tumor Challenge**

The B16-F1 murine melanoma (American Type Culture Collection, ATCC) and the B16.OVA melanoma expressing a truncated OVA protein (31) were maintained in complete Iscove’s modified Dulbecco’s medium as described (1), and extended in vitro passaging was avoided. For tumor challenge, cells were washed 3× in medium, and 104 tumor cells were injected s.c. into the flank of mice. Tumor size and survival were calculated as described (1).

**MSU + Msmeg Treatment**

Mice were treated with 2 × 104 colony-forming units (CFU) of Msmeg (mc2155) and 250 μg MSU crystals (containing <0.01 EU/10 mg) in a total volume of 100 μl PBS as described.
(9). Full treatment involved four injections of MSU + Msmeg given s.c. around the tumor, starting when the tumor became palpable (usually around days 7–9) and repeated every second day thereafter. Control mice received 100 µl PBS (Invitrogen).

**Flow Cytometry**

Lymph nodes or tumors were digested using DNase I and Liberase TL (Roche) as described (9). Blood was collected in Alsever’s solution, and red blood cells were lysed in ammonium-chloride-tris buffer (both made in house). Single cell suspensions were then resuspended in FACS buffer (PBS with 10 mM EDTA, Sigma; 2% FBS, Gibco; and 0.01% NaN₃, Sigma) and blocked with anti-mouse CD16/32 (2.4G2) before staining with fluorochrome antibodies specific for the following markers: CD45 (30F11), CD45.1 (A20), CD4 (GK1.5), CD8α (53-6.7), CD11c (HL3), CD11b (M1/70), Vo2 (B20.1), and Vβ5.1/5.2 (MR9-4), all from BD Biosciences; Ly6C (HK1.4), CD44 (X54-5/7.1), CD135 (A2F10), and MCHCII (M5/114.15.2) from Biolegend; CD115 (AFS98), CD45.2 (104), and Ly6G (1A8) from ebioscience; Ly6B (7/4) from AbD Serotec; and CCRII (475301) from R&D Systems. Anti-CD4 (GK1.5), -CD8α (2.43), and -MCHCII (3JP) were affinity purified from hybridoma supernatants. Streptavidin-PE, -APC, or PE-Texas-Red (BD Biosciences) were used where required. Dead cells were excluded by staining with 6’-diamidino-2-phenylindole (DAPI) or Live/Dead Fixable blue (Invitrogen). For intracellular cytokine staining, cell suspensions were incubated for 6 h in 1 µg/ml Golgi Stop (BD Biosciences) and 2 µg/ml Brefeldin A (eBioscience) with no restimulation. After surface staining, cells were stained with anti-IL-12p40 (C17.8 from ebioscience), anti-TNF-α (MP6-XT22, from BD Biosciences) antibodies, or isotype controls (R3-34 BD Biosciences or EBRG1, ebioscience), using the BD Cytofix/Cytoperm kit (BD Biosciences). Staining for iNOS was done in the same way with M-19 polyclonal antibody (sc-650, Santa Cruz Biotechnology) or control rabbit IgG (sc-2027, Santa Cruz Biotechnology) but without incubation with Golgi Stop/Brefeldin A. Acquisition was performed on a BD LSRII SORP or a BD LSR Fortessa SORP (Becton Dickinson), and data were analyzed using FlowJo Version 9.8.3 (Tree Star).

**DC Depletion**

For DC depletion, CD11c-DTR BM chimeras were injected i.p. with 15 ng diphtheria toxin (DT) (Sigma-Aldrich) per gram body weight 12 h prior to each MSU + Msmeg treatment. In some experiments, depletion was confirmed by flow cytometry of tumor-draining LNs and tumors as shown in Figure S1 in Supplementary Material.

**In vivo T Cell Proliferation**

CD8⁺ T cells from spleens and LNs of OTI × B6.SJL-Ptprca mice were positively selected using anti-CD8⁺ MACS Beads (Miltenyi Biotec) and labeled with CFSE as described (32). Purity was >87% as assessed by positive staining with anti-CD8α, -Vo2, and -Vβ5 antibodies. About 2 x 10⁶ cells were injected i.v. into C57BL/6J or CD11c-DTR BM chimeric hosts bearing established B16.OVA tumors. OTI proliferation was assessed in tumor-draining LNs 6 days later.

**Inhibition of CSF1 Signaling**

The small-molecule GW2580, a selective inhibitor of the CSF1 receptor kinase (33), was used to block CSF1 signaling in vivo. Tumor-bearing mice received six daily doses of either 160 mg/kg GW2580 (kindly provided by Prof Peter Shepherd, University of Auckland, New Zealand) in a solution of 0.5% methylcellulose (Sigma) and 0.1% Tween80 (Sigma) in dH₂O, or control diluent, by oral gavage starting 4 h before the first MSU + Msmeg treatment. This dose was previously shown to achieve 100% inhibition of CSF1-dependent monocyte growth (33).

**Monocyte Isolation from BM and Adoptive Transfer**

Monocytes were enriched from BM as described (15, 34) with minor modifications. BM cell suspensions were incubated with biotinylated antibodies specific for the following markers: B220 (RA3-6B2), CD19 (1D3), CD49b (DX5), CD90.2 (53-2.1), and Ly-76 (Ter119) (all from eBioscience); Ly6G (1A8) and CD135 (CSF1R). Recipients were injected s.c. around the tumor, starting when the tumor became palpable. To test whether DCs are required for therapeutic activity, we used CD11c-DTR BM chimeric hosts bearing established B16.OVA tumors. OTI proliferation was assessed in tumor-draining LNs 6 days later.

**CCL2 Blocking**

Mice bearing palpable B16.OVA tumors were injected i.p. with 100 µg anti-CCL2 antibody (clone 2H5, Biolegend) or Armenian hamster IgG control antibody (clone HTK888, Biolegend) 2 h prior to i.v. transfer of purified BM monocytes.

**Statistical Analyses**

Statistical analyses were done using GraphPad Prism 5. Two-tailed Student’s t-test was used to compare two groups. Multiple groups were compared by one-way ANOVA with Tukey’s post-test, or Kruskal-Wallis test with Dunn’s post-test for data that were not normally distributed. Survival was analyzed using the log-rank test with Bonferroni’s correction for multiple testing. Differences of p < 0.05 were deemed significant (*), p < 0.01 very significant (**), and p < 0.001 extremely significant (***)

**RESULTS**

**DCs Are Required for Tumor Immunotherapy by MSU + Msmeg**

Treatment of established B16 melanomas with the immune-activating agents MSU + Msmeg delays tumor growth and increases survival (9, 11). Treatment success requires adaptive immunity, in particular CD8⁺ T cells, and correlates with the accumulation of moDCs in dLN as early as 1 day after the first MSU + Msmeg treatment (9). To test whether DCs are required for MSU + Msmeg immunotherapy, we used CD11c-DTR BM chimeras and depleted them of CD11c⁺ cells by administration of DT throughout the duration of MSU + Msmeg treatment. Depletion of CD11c⁺ cells abrogated the treatment-induced immune response.
reduction in tumor size and increase in survival (Figures 1A,B). Successful depletion of CD11c+MHCII+ DCs in dLN after the last treatment was verified in separate experiments (Figure 1C) and confirmed that different DCs subsets, including CD8α+ DCs, CD11b+ DCs, CD8α+CD11b– DCs, and CD11b+CD64+Ly6C+ moDCs, were all significantly reduced in DT-treated CD11c-DTR chimeras (Figure 1D; Figure S1 in Supplementary Material). Thus, CD11c+ DCs are necessary for MSU + Msmeg antitumor activity.

**MSU + Msmeg Recruits Monocytes from Blood to dLN and Induces Expression of DC Markers**

We sought to establish the origin of the moDCs induced by MSU + Msmeg treatment. A substantial population of moDCs was observed in the dLN of intact mice that received MSU + Msmeg subcutaneously. MoDCs formed a distinct population within the CD11c+MHCII+ DCs, uniformly expressing high levels of CD11b, Ly6C, Ly6B, and CD64 (Figure 2A). The numbers of moDCs and monocytes in dLN were comparable to the numbers seen in similarly treated tumor-bearing animals [Figure 2B (9)] and represented about 10% of the total DC population in treated dLNs. In contrast, the proportions of other DC subsets were not significantly affected by MSU + Msmeg treatment (9).

We then examined the fate of purified BM monocytes transferred into the blood of recipient mice. Transferred cells were recruited to MSU + Msmeg-treated dLN (Figures 2C,D), and by 17 h after transfer they had already upregulated expression of the DC markers CD11c and MHCII (Figure 2E). Thus, the early influx of moDCs into the dLN of MSU + Msmeg-treated mice is likely due to monocytes entering the dLN directly from

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**FIGURE 1** | Dendritic cells are required for the activation of antitumor immunity by MSU + Msmeg. C57BL/6 WT (C57WT) or CD11c-DTR BM chimeras were injected with B16 melanoma tumors, treated every second day for four times with MSU + Msmeg or PBS, and at the same time depleted of CD11c+ cells by i.p. DT treatment as indicated. (A) Individual tumor sizes in each group, as measured on the day when average tumor size in the C57 WT control group approached 150 mm² (days 17–23 after tumor challenge). Horizontal lines show mean ± SEM. (B) Survival of tumor challenged mice. Data in (A,B) are pooled from two independent experiments each with 3–5 mice/group. (C) Numbers of total DCs (CD11c+MHCII+) and (D) numbers of moDCs (CD11c+MHCII+CD11b+CD64+Ly6C+) in draining and contralateral LNs 2 days after the completion of MSU + Msmeg treatment. Data refer to one of two independent experiments that gave similar results. Graphs show mean ± SEM for 5 mice/group. Statistical analysis in (A,C,D) was by one-way ANOVA with Tukey’s post-test. (B) Log-rank test with Bonferroni’s correction for multiple testing.
**FIGURE 2** | Monocyte accumulation and upregulation of DCs markers in the dLN of MSU + Msmeg-treated mice. (A,B) Mice were injected s.c. with MSU + Msmeg or PBS on days 0 and 2. About 17 h later dLNs were harvested and analyzed by flow cytometry. (A) Expression of DC and monocyte lineage markers on total DCs (MHCII+CD11c+), CD11b+ DC (CD11b+CD8−Ly6B−Ly6C− DCs), CD8α+ DC (CD8α+CD11b− DCs), and moDCs (CD11b+Ly6B+Ly6C+ DCs). (B) Frequency and number of moDCs (CD11b+Ly6B+Ly6C+CD11c−MHCII+) and monocytes (CD11b+Ly6B+Ly6C+CD11c+) in dLN. Data are pooled from two independent experiments each with 3–5 mice/group. (C,D,E) As in (A) but mice received two million CD45.1+ BM monocytes i.v. 17 h before analysis. (C) Identification of transferred monocytes in dLN. (D) Number of transferred monocytes and (E) expression of CD11c and MHCII on transferred monocytes, expressed as median fluorescence intensity (MFI). Data in (C–E) are from one of two independent experiments, each with 3–5 mice/group, that gave similar results. All graphs show mean + SEM. Statistical analyses used a Student’s t-test.
the blood, rather than monocytes migrating from the tumor to the dLN. However, these data do not rule out that, at later time points, moDCs in dLN may originate from either tumor or blood.

**CCR2+ Monocytes Enter dLN and Tumors in a CCL2-Dependent Manner**

Blood monocytes can be divided into two subsets: Ly6C<sup>hi</sup> patrolling monocytes that migrate in and out of tissues in the steady state, and Ly6C<sup>hi</sup> inflammatory monocytes that express CCR2 and are actively recruited to sites of inflammation (35). We therefore examined CCR2 expression on monocytes, moDCs, and cDCs in the dLN of mice treated with MSU + Msmeg 45 h earlier. Monocytes and moDCs both expressed CCR2, while other Ly6B-Ly6C<sup>-</sup> DCs were mostly CCR2 negative (Figure 3A). Consistent with this observation, the frequency of blood monocytes expressing CCR2<sup>+</sup> was diminished 15 h after MSU + Msmeg treatment, suggesting recruitment into inflamed tissues (Figure 3B).

We also investigated the role of CCL2 in the recruitment of monocytes to the dLN. CCL2, also known as MCP-1, is the main ligand of CCR2, but its role in the migration of monocytes to sites of inflammation is controversial (19, 36). Systemic treatment with CCL2-blocking antibodies almost completely inhibited the recruitment of both i.v. transferred BM monocytes and host monocytes to the dLN (Figure 3C). The proportions of transferred and host monocytes in tumors were also significantly reduced in anti-CCL2-treated mice (Figure 3D). The effects of CCL2 blockade on tumor progression were not determined. Thus, CCR2<sup>+</sup> monocytes enter the dLN and tumors of MSU + Msmeg-treated mice in a CCL2-dependent manner.

**Blocking Monocyte and moDC Recruitment Abolishes MSU + Msmeg-Induced Antitumor Activity**

To begin to address the contribution of moDCs to the antitumor immune response induced by MSU + Msmeg, we used treatment with GW2580, a small-molecule inhibitor highly specific for the CSF1R kinase (33). CSF1 promotes proliferation, survival, and differentiation of monocytes, and blocking CSF1R signaling impairs monocyte recruitment without affecting conventional DC subsets [Ref. (37) and data not shown].

Treatment with GW2580 induced a marked reduction in monocytes and moDCs in the dLNs (Figure 4A) and tumors (Figure 4B and data not shown) of MSU + Msmeg-treated mice. In addition, administration of GW2580 to prevent monocyte and moDC recruitment throughout the duration of MSU + Msmeg treatment completely abrogated the antitumor effect (Figure 4C). We therefore conclude that moDCs are required for MSU + Msmeg tumor therapy.

**Monocytes and moDCs Produce Proinflammatory Factors and Are Necessary for Proliferation and Tumor Infiltration of CD8<sup>+</sup> T Cells**

To elucidate the function of moDCs in mediating the antitumor activity of MSU + Msmeg, we chose to first investigate whether they produce proinflammatory cytokines and iNOS. During infections, moDCs are well known to be major producers of TNFα and iNOS (13) and can also be a critical source of IL-12 (15, 18). In MSU + Msmeg-treated mice, the majority of moDCs in dLN were positive for intracellular iNOS and TNFα (Figures 5A,B). In addition, about 15% of moDCs contained both intracellular TNFα and IL-12p40 (Figures 5A,B, right panel). Monocytes in the same dLN showed a similar pattern of intracellular iNOS, TNFα, and IL-12p40 staining, but the proportion of cytokine-expressing cells was lower compared to moDCs (Figure 5C).

Monocyte-derived DCs have been reported to crossprime CD8<sup>+</sup> T cells (38, 39) and induce strong Th1 and cytotoxic T lymphocyte responses (14, 15, 19). We used B16 tumors expressing the model antigen OVA together with adoptive transfer of CFSE-labeled naive CD45.1<sup>+</sup> OTI T cells to assess priming of tumor-specific CD8<sup>+</sup> T cells in response to MSU + Msmeg. Consistent with our previous work (9), MSU + Msmeg treatment increased the number of total and divided OTI cells in dLN (Figure 6A). Reducing the number of moDCs in dLN by treatment with GW2580 inhibited the increase in OTI proliferation (Figure 6A), indicating that moDCs were necessary for this response. We then examined the percentage of CD8<sup>+</sup> T cells and OTI cells in tumors. Again, GW2580 fully abrogated the increase in CD8<sup>+</sup> T cell and OTI cell infiltration into MSU + Msmeg-treated tumors (Figure 6B). These data indicate that moDCs are necessary for the efficient proliferation of tumor-specific T cells in the dLN of MSU + Msmeg-treated mice, and for their accumulation in the tumor.

**moDCs Are Sufficient for MSU + Msmeg Antitumor Activity**

To assess whether moDCs are sufficient for the development of antitumor immunity after MSU + Msmeg treatment, we used an experimental setup similar to the one outlined in Figure 7A to generate mice in which moDCs were the only DC subset present. Tumor-bearing CD11c-DTR BM chimeric mice were injected with DT to deplete CD11c<sup>+</sup> cells throughout the duration of MSU + Msmeg treatment. At the same time, mice were adoptively transferred with BM monocytes from wild-type (WT) C57BL/6 mice. As WT monocytes cannot express DTR, they were not affected by DT treatment and could differentiate into CD11c<sup>+</sup> moDC when exposed to the appropriate conditions. In some experiments, chimeras were also injected with B16.OVA tumors and adoptively transferred with naive OTI CD8<sup>+</sup> T cells to assess the response of tumor-specific CD8<sup>+</sup> T cells.

B16.OVA tumors were analyzed for percentage of total CD8<sup>+</sup> T cells and antigen-specific OTI cells at the end of MSU + Msmeg treatment. Depletion of CD11c<sup>+</sup> cells at the time of MSU + Msmeg treatment abrogated the increase in CD8<sup>+</sup> T cell infiltration into the tumor. Transfer of WT monocytes into CD11c-depleted mice was sufficient to partially recover CD8<sup>+</sup> T cell accumulation (Figure 7B, left panel) and completely restored the frequency of OTI T cells to levels comparable to those in CD11c-sufficient controls (Figure 7B, right panel). Transfer of monocytes also restored the antitumor activity of MSU + Msmeg, resulting in reduced tumor size (Figure 7C) and increased survival similar to WT controls (Figure 7D). Recovery of the antitumor response
Mice. As WT monocytes cannot express DTR, they were not transferred with BM monocytes from wild-type (WT) C57BL/6. Tumor-specific CD8+ was sufficient to partially recover CD8+ T cells in the tumor. Transfer of WT monocytes into CD11c-depleted mice MSU+ was able to abrogate the increase in CD8+ T cell infiltration into MSU tumors. CFSE-labeled naive CD45.1+ OTI T cells to assess priming of OTI T cells and antigen-specific OTI cells at the end of MSU+ treatment. Depletion of CD11c+ cells throughout the duration of treatment increased the number of total and divided OTI cells in dLN (Figure 6A). Reducing the number of moDCs in dLN by treatment with GW2580 inhibited the increase in OTI cell frequency. In some experiments, chimeras were also injected with B16.OV A tumors to generate mice in which moDCs were the only DC subset present.

Figure 5C). In MSU+ treated tumors (Figure 6B), the majority of moDCs in dLN showed a similar pattern of intracellular iNOS, TNFα were positive for intracellular iNOS and TNFα were positive for intracellular iNOS. During MSU+ treatment, we used an Msmeg+ expressing cells was lower compared to moDCs (Figures 5A,B). In MSU+ treated mice, the majority of moDCs in dLN -treated mice, the majority of moDCs in dLN. Monocyte-derived DCs have been reported to crossprime T cells and OTI cells in tumors. Again, GW2580 fully restored the antitumor activity of MSU+ treated mice, the majority of moDCs in dLN. At the same time, mice were adoptively transferred (CD45.1+) or host (CD45.2+) monocytes (Ly6B+Ly6C−MHCII+) or other DCs (CD11c+MHCIILy6B−Ly6C+) in dLNs of mice treated with MSU+Msmeg once 45 h earlier. (B) Frequency of CCR2+ monocytes in blood 15 h after MSU+Msmeg treatment. (C,D) Mice bearing palpable B16.OVA tumors were treated with MSU+Msmeg. The recruitment of transferred (CD45.1+) or host (CD45.2+) monocytes (Ly6B+Ly6C−MHCII+CD11c+) into LN (C) and tumors (D) was examined 18 h later. Data are pooled from two independent experiments each with 3–5 mice/group. All graphs show mean ± SEM. Statistical analyses used a Student’s t-test (B) or a Kruskal–Wallis with Dunn’s post-test (C,D).

**FIGURE 3 | CCL2 is necessary for the recruitment of monocytes to MSU+Msmeg-treated dLNs and tumors. (A) Expression of CCR2 on monocytes (CD11b+Ly6B+Ly6C+CD11c+), moDCs (CD11c+MHCIILy6B+Ly6C+), and other DCs (CD11c+MHCIILy6B−Ly6C−) in dLNs of mice treated with MSU+Msmeg once 45 h earlier. (B) Frequency of CCR2+ monocytes in blood 15 h after MSU+Msmeg treatment. (C,D) Mice bearing palpable B16.OVA tumors were treated with MSU+Msmeg. The recruitment of transferred (CD45.1+) or host (CD45.2+) monocytes (Ly6B+Ly6C+CD11c+) into LN (C) and tumors (D) was examined 18 h later. Data are pooled from two independent experiments each with 3–5 mice/group. All graphs show mean ± SEM. Statistical analyses used a Student’s t-test (B) or a Kruskal–Wallis with Dunn’s post-test (C,D).**
did not require transfer of OTI T cells and occurred regardless of challenge with B16 or B16.OVA tumors, or OTI transfer. These results indicate that, in a situation where other CD11c+ cells are depleted, moDCs are sufficient for induction of tumor-specific CD8+ T cell responses and successful tumor immunity in response to MSU + Msmeg.

**DISCUSSION**

In this study, we investigated the role of moDCs in the antitumor immune response elicited by local treatment with MSU + Msmeg. We show that DCs are necessary for the activity of MSU + Msmeg, as depletion of CD11c+ cells completely abrogated the antitumor effect. Interestingly, blockade of CSF1R to selectively deplete moDCs without affecting other DC subsets also abrogated the effect of immunotherapy. In a reciprocal fashion, adoptive transfer of monocytes into mice depleted of CD11c+ cells was sufficient to restore antitumor activity. MoDCs originated from circulating monocytes in response to MSU + Msmeg treatment, expressed iNOS, TNFα, and IL-12 and were necessary for the priming of tumor-specific CD8+ T cells. In addition, moDCs were necessary and sufficient for the accumulation of tumor-specific CD8+ T cells in tumors. Together, these data show that moDCs play an essential role during immunotherapy with MSU + Msmeg.

The experiments in this paper investigated the role of moDCs using a model of B16 melanoma and immunotherapy with MSU + Msmeg. Our previous work compared several different treatments with immunostimulatory agents, including LPS, polyI:C (9), and CpG immunostimulatory oligonucleotides

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**FIGURE 4** | Treatment with the CSF1-signaling inhibitor GW2580 impairs monocyte and moDCs recruitment and abrogates the antitumor activity of MSU + Msmeg. Mice bearing B16.OVA tumors were treated with MSU + Msmeg, and at the same time they received daily oral doses of GW2580 or vehicle control. LN and tumors were analyzed 2 days after the last MSU + Msmeg treatment. (A) Numbers of monocytes (CD11b+Ly6B+Ly6C−CD11c−MHCII+) and moDCs (CD11b+Ly6B+Ly6C+CD11c+MHCII+) in dLN. (B) Frequencies of monocytes (CD45−CD11b+Ly6C+Ly6G−CD11c+), total DCs (CD45+CD11c+MHCII+), and moDCs (CD45+CD11c+MHCII+CD11b+Ly6C+Ly6B+) in tumors. (C) Tumor sizes on day 13. All data are pooled from two independent experiments each with 5 mice/group. Graphs show mean ± SEM. Statistical analyses used ANOVA with Tukey’s post-test (A) or a Kruskal–Wallis with Dunn’s post-test (B,C).
Monocytes and moDCs in dLN express proinflammatory mediators after MSU + Msmeg treatment. Mice were treated with a single dose of MSU + Msmeg s.c., and 19 h later dLNs were collected and analyzed by intracellular staining for the indicated markers. Staining controls included an isotype control (isotype) and stained samples that were not treated with GolgiStop/Brefeldin A (no block). (A) Representative flow plots and (B) frequencies of iNOS-, IL-12-, and TNFα-expressing moDCs (CD11c+MHCII+CD11b+Ly6B+Ly6C+). (C) Frequencies of monocytes (CD11b+Ly6B+Ly6C+CD11c−) expressing proinflammatory molecules. Data are from one of two independent experiments, each with 3–5 mice/group that gave similar results. Graphs show means + SEM. Statistical analyses used a Student’s t-test.
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Kuhn et al.

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igure 6 | Monocyte-derived DCs are necessary for the proliferation of tumor-specific CD8\(^+\) T cells in dLN and their infiltration into tumors. Mice bearing B16.OVA tumors were treated with MSU + Msmeg, and at the same time they were given daily oral doses of GW2580 or vehicle. LN and tumors were analyzed 2 days after the last MSU + Msmeg treatment. (A) Numbers of total and divided OTI T cells (CD45.1\(^+\)CD8\(^+\) lymphocytes) in tumor dLN, as assessed by CFSE dilution. (B) Total CD8\(^+\) and OTI T cells (CD45\(^+\)CD8\(^+\) and CD45\(^+\)CD8\(^+\)CD45.1\(^+\), respectively) expressed as frequencies of total live cells in tumors. Data are pooled from two independent experiments, each with 5 mice/group. All graphs show mean \(\pm\) SEM. Statistical analyses were by one-way ANOVA with Tukey’s post-test (A) and Kruskal–Wallis test with Dunn’s post-test (B).

(unpublished data), to show that antitumor activity correlated with increased numbers of moDCs in the tumor dLN. In addition, treatments that were effective on B16 melanoma were also effective on EL4 thymomas (11) and 4T1 mammary carcinomas (9), suggesting that the requirement for moDCs in antitumor immune responses extends to other forms of immunotherapy besides MSU + Msmeg, and to multiple tumor models.

Our finding that CCR2\(^+\)Ly6C\(^{hi}\) monocytes were recruited to MSU + Msmeg dLNs and tumors is consistent with literature showing that these monocytes are preferentially attracted to sites of inflammation (35). Accordingly, the strong reduction in monocyte recruitment after treatment with CCL2-blocking antibodies suggests that CCR2–CCL2 interactions are largely responsible for the attraction of monocytes to the site of MSU + Msmeg treatment (23, 30, 36). While there is clear evidence that monocytes can differentiate into moDCs during inflammation (13, 14, 35) surprisingly little is known about the signals that govern this process. In line with our findings, CSF1 is the major lineage regulator of mononuclear phagocytes in the BM and thus is required for the development of monocytes and subsequently moDCs (40). The upregulation of MHCII and production of IL-12p40 by moDCs instead were shown to depend on natural killer (NK) cell-derived IFN\(\gamma\) in a study of Toxoplasma gondii infection (18). In our model, depletion of NK1.1\(^+\) cells did not reduce the number of MHCII\(^+\) moDCs in dLN after MSU + Msmeg treatment (data not shown), suggesting that NK cells were not vital for moDC differentiation. However, NK cells may be important for the function of moDCs, a possibility that was not examined in our experiments.

In our study, a significant proportion of moDCs expressed IL-12p40. The critical role of IL-12 for the induction of cytotoxic T lymphocytes and for antitumor immunity is well established. In addition, moDCs produced TNF\(\alpha\) and iNOS. This activity was previously used as a defining feature of monocyte-derived “TIP” DCs (13). Mounting evidence suggests that TNF\(\alpha\) is important for antitumor immunity, as shown by experiments where tumor immunotherapy with polyI:C required TNF\(\alpha\) for treatment success (41). Furthermore, a recent study showed that the combined action of TNF\(\alpha\) and IFN\(\gamma\) is necessary for the control of a range of murine and human tumors by driving cancer cells into senescence (20).

In contrast to IL-12 and TNF\(\alpha\), the role of NO/iNOS in tumors is more controversial. A number of studies have shown that macrophage- or MDSC-derived NO can suppress antitumor T cell responses in vitro [reviewed in Ref. (42)]. However, others report that NO is required for direct tumoricidal activity of macrophages in vitro (43, 44). In vivo data are unfortunately limited and suggest that the effects of NO in vivo may not always be accurately reflected in ex vivo suppression experiments (45). This notion is further supported by the finding that in a spontaneous melanoma model ROS-producing monocytic effector cells have direct antitumor activity that is independent of T cells (27). It is possible that in our model monocytes and moDCs exerted some direct antitumor activity via iNOS and TNF\(\alpha\), as these factors
Monocyte-Derived Dendritic Cells in Tumor Immunotherapy

Figure 7 | Monocyte-derived DCs are sufficient for the recruitment of CD8+ T cells to tumors and the activation of antitumor immunity. (A) Schematic representation of experimental design. C57BL/6 WT (C57WT) or CD11c-DTR BM chimeras were injected with B16 melanoma tumors, treated every second day for four times with MSU+Msmeg or PBS, and at the same time depleted of CD11c+ cells by DT treatment as indicated. Some of these mice were also injected with WT BM monocytes to obtain mice where moDC populations could only originate from the injected monocytes. (B) Mice were treated as in (A), except that tumors were B16.OVA, and all mice also received naive OTI cells given at the time of the second dose of immunotherapy. Graphs show the percentage of CD8+ T cells (CD45+CD4−CD8+T) and OTI cells (CD45+CD4−CD8+TIDR.1) among total live cells in tumors 2 days after the fourth MSU+Msmeg treatment. Data are pooled from two independent experiments each with 5 mice/group. Graphs show mean ± SEM. Statistical analysis used a Kruskal–Wallis test with Dunn’s post-test. (C) Mean tumor sizes ± SEM on day 21 after tumor challenge. Data are pooled from three independent experiments each with 5 mice/group. Statistical analysis was by ANOVA with Tukey’s post-test. (D) Survival data, pooled from two independent experiments each with 5 mice/group. Statistical significance was evaluated using a log-rank test and Bonferroni’s correction.

were expressed by monocytes and moDC in the dLN and also in the tumor (data not shown). However, MSU + Msmeg therapy of B16 melanoma was CD8+ T cell dependent (9), and CD8+ T cell proliferation was strongly diminished when moDCs were blocked. Therefore, one major role of moDCs was in supporting the priming of CD8+ T cells, although we cannot rule out additional direct antitumor functions.

We have previously shown that in the steady state CD11c+ DCs are necessary for crosspresentation to tumor-specific CD8+ T cells in dLN, as CD8+ T cell proliferation was abrogated in mice depleted of CD11c+ cells (9). The experiments presented here reveal that, during immunotherapy, blockade of CSF1R-dependent monocytes and moDCs reduced CD8+ T cell proliferation in dLN to a level similar to that observed in the steady state. Therefore, promoting the activation of adaptive antitumor immunity is likely to be a key function of moDCs during MSU+Msmeg therapy. We also observed a striking impact of moDCs on the infiltration of CD8+ T cells into tumors. This may merely reflect increased

Frontiers in Immunology | www.frontiersin.org
11 November 2015 | Volume 6 | Article 584
they promote tumor growth (21, 54). Even tumor-infiltrating
associate with increased survival in patients (53), but more often
nate tumors through secretion of cytotoxic factors (44, 45) and be
breast cancer (52). In a similar manner, macrophages can elimi-
cate that monocytic cells may be important for antitumor responses
variable results of CSF1/CSF1R blockade, CSF1 gene signatures
tumor-specific T cells during immunotherapy. In line with the
difference with CSF1 inhibition alone (46, 48, 49), or even suggest
detrimental effect on tumor therapy, not only by depleting mac-
(27, 29, 50). Here, we show that CSF1R inhibition can have a
that monocytic cells may be important for antitumor responses
(27, 29, 50). Here, we show that CSF1R inhibition can have a
detrimental effect on tumor therapy, not only by depleting mac-
expression of receptors to sense the local environment
inflammatory environment (9, 29). Furthermore, therapies may also preferentially activate
DCs depending on the expression of recep-
tors for damage- and pathogen-associated molecular patterns in
these cell populations. A systematic study comparing the effect
of different treatments on several DC subsets and in a range of
tumors would be needed to fully address this issue.
CSF1-dependent monocytes and monocyte-derived cells
including tumor-associated macrophages, MDSCs, and moDCs
express a vast range of receptors to sense the local environment
and are highly plastic in their response to various stimuli. This may
explain why several studies show improved antitumor responses
after CSF1/CSF1R blockade when combined with radiotherapy,
chemotherapy, or antiangiogenic therapy (21, 37, 46–48) but no
difference with CSF1 inhibition alone (46, 48, 49), or even suggest
that monocytic cells may be important for antitumor responses
(27, 29, 50). Here, we show that CSF1R inhibition can have a
detrimental effect on tumor therapy, not only by depleting mac-
rophage effector cells (51) but also by impairing the activation of
tumor-specific T cells during immunotherapy. In line with the
variable results of CSF1/CSF1R blockade, CSF1 gene signatures
themselves were found to be either associated with poor survival
or with improved survival, depending on the specific subtype of
breast cancer (52). In a similar manner, macrophages can elimi-
nate tumors through secretion of cytotoxic factors (44, 45) and be
associated with increased survival in patients (53), but more often
they promote tumor growth (21, 54). Even tumor-infiltrating
MDSCs, generally thought to be highly immune suppressive,
have been shown to exhibit plasticity and acquire the ability to
crossprime antitumor CD8+ T cells in response to IL-12 (55).
CSF1R inhibitors are currently being tested in clinical trials
cancer therapy in patients. However, our data indicate that
in some cases CSF1 may actually be critical for the initiation
antitumor responses. This highlights the need for caution in
combining CSF1R blockade with other therapies. The role of the
immune system in mediating antitumor responses not only after
immunotherapy but also in certain chemotherapies and radio-
therapy is increasingly recognized. Therefore, it is important to
fully delineate the role of monocytic cell populations in different
types of tumors and treatments before carefully considering the
situations in which a combination with CSF1 blockade may be
beneficial. Our results indicate that, for the purpose of tumor
therapy, blocking or eliminating monocytic cells will likely be less
productive than appropriately activating them to harness their
properties and promote powerful antitumor immune responses.

AUTHOR CONTRIBUTIONS
SK and FR conceived the study; SK and JY executed experiments;
SK, JY, and FR designed and analyzed experiments; SK and FR
wrote the paper; SK, JY, and FR commented on the manuscript
and approved it for submission.

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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