CX3CR1+ CD115+ CD135+ common macrophage/DC precursors and the role of CX3CR1 in their response to inflammation.

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Monocytes, macrophages, and DCs form networks of phagocytic cells throughout most tissues, the development of which are dependent on the CSF-1 receptor (cSF1r, also known as CD115, c-fms, and M-CSF receptor) (1,2). These cells, which are sometimes referred to as the mononuclear phagocyte system, play major roles in development, scavenging, inflammation, and antipathogen defenses (3,4). They are highly heterogeneous in phenotype, tissue distribution, and function (3,5,6). Considerable attention is currently focused on the characterization of their progenitors and precursors, the signals driving their development in the BM, their migration to tissues, and their homeostasis in peripheral tissues. CSF-1R and its two known ligands, M-CSF and IL34 (7), are critical for the development of this lineage because M-CSF-deficient mice (op/op and cSF1r−/−) have a milder phenotype than the Csf1r-deficient mice (8). Other cytokines, such as GM-CSF, FLT3, LT-α1β2 (LT-α) (9–15), and chemokines (16,17) have also been shown to control the development and homeostasis of the macrophage and DC networks. Cellular cloning and transplantation studies have shown that many macrophage subsets, most of the conventional DCs (cDCs) in the secondary lymphoid organs of mice, and at least a fraction of the DCs in the mouse thymus probably originate from myeloid progenitors (18–20).

CX3CR1 expression is associated with the commitment of CSF-1R+ myeloid precursors to the macrophage/dendritic cell (DC) lineage. However, the relationship of the CSF-1R+CX3CR1+ macrophage/DC precursor (MDP) with other DC precursors and the role of CX3CR1 in macrophage and DC development remain unclear. We show that MDPs give rise to conventional DCs (cDCs), plasmacytoid DCs (PDCs), and monocytes, including Gr1+ inflammatory monocytes that differentiate into TipDCs during infection. CX3CR1 deficiency selectively impairs the recruitment of blood Gr1+ monocytes in the spleen after transfer and during acute Listeria monocytogenes infection but does not affect the development of monocytes, cDCs, and PDCs.

CX3CR1 is a G-protein-coupled receptor that is expressed on a subset of monocytes, macrophages, and DCs. It is required for the development and function of these cells, and its expression is regulated by a variety of cytokines and chemokines. The role of CX3CR1 in inflammation and immunity is underscored by the fact that mice lacking CX3CR1 are more susceptible to certain infections and inflammatory diseases. CX3CR1-deficient mice are also more susceptible to certain forms of cancer, suggesting that CX3CR1 may play a role in immune surveillance and tumor rejection.

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Granulocyte-macrophage progenitors (GMPs [reference 21]) include a clonogenic BM macrophage/DC precursor (MDP) that gives rise to spleen cDCs (both the CD11c+CD8α+CD11b− and CD11c+CD8α−CD11b+ subsets) directly, with no monocytic intermediate, and to monocytes and macrophages (9, 22, 23). The MDP has no significant granulocytic potential, and initial studies failed to detect a plasmacytoid DC (PDC) potential (9, 22). Another precursor, common DC precursor (CDP), was recently shown to give rise to cDCs and PDCs but not to monocytes, and it did not respond to CSF-1 (24, 25). This result was interpreted as indicating the existence of two pathways for DC generation. However, MDPs and CDPs are both included in the CD115+lin− fraction of BM progenitors (9) and could represent different stages of differentiation along the same pathway. It is also possible that differences in differentiation potential between these cells reported by different groups may reflect differences in experimental protocols rather than intrinsic properties of the cells.

The chemokine receptor and adhesion molecule CX3CR1 is not expressed on early hematopoietic progenitors and is first detected on MDPs. CX3CR1 is therefore associated with the commitment of myeloid progenitors to the monocyte/macrophage/DC lineage (22). However, its role in the development and homeostasis of cells of the mononuclear phagocyte system remains unknown.

In this paper, we therefore reevaluated the differentiation potential of the MDP and the possible roles of CX3CR1 in the differentiation of mononuclear phagocytes in mice using adoptive transfer and disease models. We found that MDPs can give rise to PDCs, as well as to cDCs and monocytes, after adoptive transfer and that MDPs and CDPs share a similar surface phenotype (Lin−IL7Ra−CD117intCD135+CD115+CX3CR1+). The use of AFS98, an antibody designed to block CSF-1 binding to its receptor CD115, and CSF-1−dependent proliferation (26–28) to purify MDP did not impair the ability of MDP to give rise to monocytes, cDCs, or PDCs in vivo. Because MDP can give rise to PDCs, cDCs, and monocytes/macrophages, whereas CDP only give rise to PDCs and cDCs (24, 25), MDP appears to exhibit a broader differentiation potential than CDP and may represent an earlier precursor. CX3CR1 deficiency decreased the recruitment into the spleen of CD115+Gr1+ monocytes (TipDC precursors) after irradiation and during acute Listeria monocytogenes infection and decreased the efficiency of bacterial clearance but did not affect the development of cDCs or PDCs. The results from this study, therefore, clarify the family tree of mononuclear phagocytes and uncover the role of CX3CR1 in Gr1+ monocyte recruitment to the spleen during inflammation and infection.

RESULTS

MDPs and CDPs are phenotypically overlapping cell populations in mouse BM

Expression of the chemokine receptor CX3CR1 in GMPs (Lineage−CD117+Scal−IL7Rα−CD34+CD16/32+BM cells) (21) characterizes the MDP (22) and is thus associated with the commitment of myeloid progenitors toward the macrophage/DC lineage. The MDP is also characterized by a low expression of CD117 (c-kit, the receptor for stem cell factor), as compared with GMPs and CMPs (22), and expression of functional CSF-1R (CD115) and FLT3 (CD135) (9, 22). More recently, a precursor common to cDCs and PDCs (CDP) was reported and proposed to be distinct from the MDPs because the CDP gave rise to both cDCs and PDCs but not to monocytes/macrophages, whereas the MDPs give rise to cDCs and monocytes/macrophages but not to PDCs (24, 25). We performed an analysis of mouse BM Lin+ precursors by flow cytometry (Fig. 1 a), and the results indicated that most CDPs expressed CX3CR1 at levels similar to its level of expression on MDPs, and that most MDPs expressed both CSF-1R/CD115 and CD135 at levels similar to their expression on CDPs (Fig. 1). These data confirm the data from Waskow et al. (9) suggesting that MDPs and CDPs had an overlapping phenotype in the BM, and we therefore sought to reevaluate the differentiation potential of MDPs.

MDPs give rise to monocytes, cDCs, and PDCs

Initial studies failed to identify PDCs in the progeny of MDPs after in vivo transfer into C57BL/6 mice (9, 22, 23). However, MDPs in spleen express a CSF-1R−EGFP transgene (2), and we observed that PDCs also expressed CX3CR1 (Fig. S1, available at http://www.jem.org/cgi/content/full/jem.20081385/DC1) and that MDP gave rise to a population of CD11b−CD11c+CX3CR1+splenoocytes that are distinct from cDCs and monocyte/macrophages and may correspond to PDCs after in vivo transfer into irradiated hosts (Fig. S1). In addition, the availability of PDCA1 antibody now permits a better phenotypic definition of PDC (24, 25). We thus investigated the differentiation potential in vivo of MDP, purified as previously described (Fig. S2) (22), after i.v. adoptive transfer. BM MDP from CX3CR1+CD11b+ mice of the Cd45.1/Cd45.2 genotypes was injected i.v. into Cd45.2-irradiated hosts, and splenoocytes were analyzed by flow cytometry 6–7 d after transfer. Results indicated that MDP gave rise to donor-derived PDCA1+CD11c+CD11b−CX3CR1+PDC, as well as to CD11c+CD11b+CX3CR1− and CD11c+CD11b+CX3CR1+ cDC and to CD11c+CD11c−CX3CR1+ monocytes (Fig. 2, a–c). As expected, the frequency of donor-derived PDCs and monocytes was 2–3-fold lower than that of CD11b−CD11c− cDCs (Fig. 2 c). However, CD11b− PDCs were easily distinguishable from CD11b− cDCs by their expression of CX3CR1 and PDCA1 and their low expression of CD11c (Fig. 2, a and b). MDP-derived CD11b+ CD11c− monocytes expressed high levels of CX3CR1 (Fig. 2 a), which unambiguously distinguished them from granulocytes which do not express CX3CR1 (29).

Because the anti-CD115 antibody (AFS98) used to purify CDP (25), and MDP in one study (9), was originally screened for its ability to block the binding of CSF-1 to its receptor (26–28, 30, 31), we investigated whether labeling of MDPs with AFS98 would affect the differentiation of MDPs in our experimental model. We observed that, in vitro, the cloning...
efficiency of single MDP seeded into 96-well plates in the presence of CSF-1, as well as the size of colonies, was reduced when AFS98 was added to the antibody cocktail used for cell sorting (Fig. S3, available at http://www.jem.org/cgi/content/full/jem.20081385/DC1). However, addition of AFS98 to the antibody cocktail used for cell sorting did not affect the differentiation potential of MDP in vivo (Fig. 2 c).

These data indicate that MDP and CDP share expression of CX₃CR1, CD115, and FLT3 and have an otherwise overlapping phenotype in the BM, whereas MDP appears to have a broader differentiation potential than CDP after adoptive transfer in vivo because MDP gave rise to both DCs (PDCs and cDCs) and monocyte/macrophages. It is of note that addition of AFS98 to the antibody cocktail used for cell sorting decreased the proliferative response of MDP to CSF-1 in vitro but did not affect their differentiation into to DCs or monocytes in vivo.

**CX₃CR1-deficient MDPs have a decreased potential to give rise to spleen monocytes**

Because expression of CX₃CR1 is associated with the commitment of myeloid progenitors to the monocyte/macrophage/DC lineage, we investigated the role of CX₃CR1 in the homeostasis of this lineage. The number of MDPs in the BM was not affected by CX₃CR1 deficiency (Fig. 3 a) and, on average, 6.5 × 10⁴ MDPs were recovered per femur from CX₃CR1⁺⁻ and CX₃CR1⁻⁻⁻ mice, suggesting that CX₃CR1 is dispensable for MDP development in the BM. We thus investigated the role of CX₃CR1 in DC and monocyte development from MDP by studying the fate of MDP in competitive adoptive transfer, in which 10⁴ MDP from each of two different donors of the Cd45.2 and Cd45.2/Cd45.1 genotype were coinjected i.v. into a Cd45.1 congenic recipient (Fig. 3 b). When both donor mice were of the CX₃CR1⁺⁻⁻ genotype, CD45.2 and CD45.2/CD45.1 MDPs contributed equally to cDCs, PDCs, and CD11b⁺ CD11c⁻ monocytes (Fig. 3, b and c). When CD45.2 MDPs were of the CX₃CR1⁻⁻⁻ genotype and CD45.2/CD45.1 MDPs were of the CX₃CR1⁺⁻⁻ genotype, they contributed equally to spleen DCs and PDCs (Fig. 3, b and d). However, CX₃CR1-deficient MDP generated spleen CD11b⁺ CD11c⁻ cells with an efficiency of one fifth to one tenth that of the control (Fig. 3, b and d).

MDP-derived CD11b⁺ CD11c⁻ splenocytes were negative for NK1.1, CD3, CD19, and Ly6G and expressed high levels of CX₃CR1 and intermediate levels of F4/80, and most of these cells expressed Gr1 (Fig. 3 e). These characteristics are similar to those of CD115⁺ Gr1⁺ blood monocytes (29) and of spleen monocytes, as recently described by Nahrendorf et al. (32). Thus, we concluded that CX₃CR1 may be selectively involved in the development, recruitment, proliferation, and/or survival of CD11b⁺ CD11c⁻ monocytes in the spleen.

**CX₃CR1 is important for the recruitment of blood Gr1⁺ monocytes to the spleen during infection and for the clearance of Lm in mice**

The only known ligand of CX₃CR1 is the transmembrane chemokine fractalkine/CX₃CL1, which is expressed in neurons, endothelial cells, and DCs (33–37). CX₃CR1 is involved in the adhesion of leukocytes, including monocytes in particular, to endothelial cells (38–42) and in the migration of microglial cells (43). Fractalkine is expressed in the T cell areas of lymph nodes (37). PCR and in situ hybridization experiments indicated that CX₃CL1 was also expressed in the spleen (Fig. 4, a and b; and Fig. S4, available at http://www.jem.org/cgi/content/full/jem.20081385/DC1). Spleen CX₃CL1-producing cells were located at the periphery of the B cell follicle.

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**Figure 1. MDP and CDP share a similar phenotype.** BM cells from CX₃CR1⁺⁺⁺⁺ reporter mice were labeled with antibodies against lineage markers (CD11c, CD11b, NK.1.1, CD3, Ter119, CD19, and Gr1), ckit (CD117), FLT3 (CD135), and CSF-1-R (CD115) and analyzed by FACS. CDP and MDP are overlapping populations; lineage⁻⁻⁻⁻⁻ CD117⁻²⁵⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻⁻˓
which correspond to the marginal zone/T cell area (Fig. 4, a and b). The precise identification of CX3CL1-expressing cells in the spleen will require the availability of specific antibodies; however, localization of Cx3cl1 messenger RNA is compatible with the expression of CX3CL1 in the marginal zone/T cell area of the spleen and, thus, with a role of CX3CL1 in the recruitment of blood monocytes.

We tested the role of CX3CR1 for the recruitment of monocytes in the spleen by adoptive transfer of monocytes and during infection with Lm. Short-term competitive transfer of BM monocytes from donors of Cd45.2 and Cd45.2/Cd45.1 genotype into irradiated Cd45.1 congenic recipients indicated that CX3CR1-deficient monocytes were only one tenth as efficient as control monocytes at accumulating in the spleen (Fig. 4 c). During Lm infection in CX3CR1-deficient BALB/c mice, the number of monocytes was increased in the blood, whereas the number of monocytes in the spleen was decreased in comparison with controls (Fig. 4 d). A similar phenomenon was observed in C57BL/6 mice (Fig. 5, e and f) both for high ($3 \times 10^5$ Lm; Fig. 4 e) and low ($7 \times 10^3$ Lm; Fig. 5 f) numbers of bacteria. To investigate whether the proliferation of blood monocytes in the spleen of Lm-infected mice was involved in their accumulation in the spleen, monocytes from the BM and spleen of infected mice at 16 and 48 h after infection were isolated by flow cytometry and analyzed for DNA content using propidium iodide (PI) staining (Fig. 4 g and Fig. S5, available at http://www.jem.org/cgi/content/full/jem.20081385/DC1). Results indicated that BM monocytes, but not spleen monocytes, actively proliferate at 16 and 48 h.

Figure 2. Differentiation potential of MDP in vivo. (a–c) MDPs from Cd45.1/2 or Cd45.2 Cd3.2 reporter mice were purified as described in Fig. S2 (available at http://www.jem.org/cgi/content/full/jem.20081385/DC1) and were adoptively transferred into irradiated (900 rad) C57BL/6 Cd45.2 congenic recipients. Spleens of recipient mice were analyzed at day 7 after transfer by flow cytometry, using lineage marker (NK1.1 CD3 CD19), CD11b, CD11c, and CD8-α antibodies. R1 corresponds to PDCs (Lin-CD11cint CD11b- CX3CR1+ cells), R2 and R3 correspond to cDCs (Lin-CD11chigh CD11b- CX3CR1+ cells and Lin-CD11cint CD11b+ CX3CR1+ cells), and R4 corresponds to monocytes (Lin-CD11b+ CD11c- CX3CR1+ cells). The experiment was repeated five times with two to three mice per group and with similar results. (b) Donor-derived Lin-CD11cint CD11b- CX3CR1+ cells express PDCA1. (c) Role of anti-CD115 Ab. The number of donor-derived cells per spleen are represented after adoptive transfer of MDP, purified with or without CD115 antibody (n = at least 3 mice per group from two experiments). Error bars show SD. (d) The flow diagram represent the Lin-CD117int CD115+ CD135+ CX3CR1+ MDPs that give rise to monocytes, cDCs, and PDCs and their putative relationship with other myeloid precursors.
CD115⁺ Gr1⁺ monocytes recruited to the spleen during *Lm* infection differentiate into effector cells (TipDCs) that produce TNF, inducible nitric oxide synthase (iNOS), and reactive oxygen intermediates (ROIs)

Because the decreased recruitment of monocytes in the spleen during *Lm* infection in CX3CR1-deficient mice correlated with a decreased bacterial clearance, we investigated whether monocytes recruited to the spleen may be involved in the control of *Lm* growth and express effector activities important for *Lm* clearance such as TNF-α secretion, iNOS expression, and reactive oxygen production. Monocytes have already been proposed to be precursors of TipDCs that accumulate to the spleen of mice infected with *Lm*, produce TNF-α and iNOS, and are needed for the control of primary infection (44–46). Recruited monocytes were the majority (90%) of the CD115⁺ Gr1⁺ subset (Fig. S7, available at http://www.jem.org/cgi/content/full/jem.20081385/DC1). In spleen, lineage⁻/Ly6G, NK1.1, CD3, and CD19) CD11b⁺ CD11c⁻ splenocytes were identified as monocytes based on their expression of CD115, Ly6C/Gr1, and F4/80 (Fig. 5a) (32).

We observed that these spleen CD115⁺ Gr1⁺ monocytes produced TNF-α, iNOS, and ROI in mice during infection after infection and, thus, that accumulation of spleen monocytes during infection is not a result of their proliferation in the spleen. In addition, the proportion of splenic monocytes that bind annexin-V in *Lm*-infected mice was similar in CX3CR1-deficient animals and controls (Fig. S6), indicating that the apoptosis rate of CX3CR1-deficient Gr1⁺ monocytes during *Lm* infection was not increased in comparison with controls.

Early control of *Lm* growth in the spleen was also significantly less efficient in CX3CR1-deficient mice on both BALB/c and C57BL/6 backgrounds (Fig. 4, h and i). Bacterial load in the spleen 24 h after an i.v. infection was 4× higher in BALB/c CX3cr1⁻/⁻ mice in comparison with controls (Fig. 5h) and at least twice as high in C57BL/6 (Fig. 4i).

Altogether, these data indicate that CX3CR1 is important for the recruitment of monocytes to the spleen in irradiated host and during *Lm* infection, and for the early control of bacteria growth. It is of note that the frequency of monocytes in the spleen of noninfected mice were not affected by CX3CR1 deficiency (Fig. 4d), indicating that CX3CR1 is dispensable for the recruitment of monocytes in the spleen in the steady state but important during acute inflammation.

Figure 3. CX3CR1 is important for the development of CD11b⁺ CD11c⁻ monocytes. (a) MDP numbers in BM from CX3CR1⁺/⁻ (black) and CX3CR1⁻/⁻ (white) mice. Data are the mean ± SD of five mice per group. (b–d) Competitive adoptive transfer of MDP into irradiated host (900 rad). Equal numbers (10⁴) of MDPs from Cd45.1/2 mice (blue) and Cd45.2/2 mice (red) were mixed and adoptively transferred into a Cd45.1 congenic recipient. Spleens of recipient mice were analyzed at day 7 after transfer by flow cytometry, using lineage marker (NK1.1 CD3 CD19), CD11b, CD11c, and CD8⁻ antibodies. (b and c) When both Cd45.1/2 and Cd45.2/2 donor mice were of the CX3CR1⁺/⁻ genotype, both donors contributed equally to CD11c⁺ CD11b⁻ (R1, cDC) and NK1.1⁻ CD3⁻ CD19⁻ CD11c⁺ CD11b⁺ (R2, cDC), NK1.1⁻ CD3⁻ CD19⁻ CD11c⁺ CD11b⁺ (R3, PDC), and NK1.1⁻ CD3⁻ CD19⁻ CD11c⁻ splenocytes (R4). In contrast, when Cd45.1/2 donor mice were of the CX3CR1⁻/⁻ genotype and Cd45.2/2 donor mice were of the CX3CR1⁺/⁻ genotype (CX3CR1 deficient), both donors contributed equally to CD11c⁻ cDCs in R1 and R2 but CD45.1/2/ CX3CR1⁻/⁻ MDPs were 10x more efficient than CD45.2/2 CX3CR1-deficient MDPs in generating CD11b⁺ CD11c⁻ splenocytes in R3. Results in b are from one representative experiment out of three, with two to three mice per experimental group, and bar graphs in c and d represent the mean and SD from three independent experiments. The asterisk indicates a significant difference between groups (P < 0.05 using the Wilcoxon test).
with *Lm* (Fig. 5, b–d; and Fig. S8) and were localized in the T cell area and in the perifollicular area of the spleen (Fig. 5 e). Altogether, these data therefore indicate that MDPs give rise to blood CD115⁺ Gr1⁺ monocytes, which are recruited to the spleen during infection via a process that involves CX₃CR1, and that these cells expressed effector functions that are important for *Lm* elimination.

Gr1⁺ monocyte survival and MDP survival and proliferation are normal in CX₃CR1-deficient mice

Other mechanisms that might be responsible for the impaired accumulation of CX₃CR1-deficient CD115⁺ Gr1⁺ monocytes in the spleen include decreased survival or proliferation of monocytes or decreased survival, proliferation, or differentiation of MDP. As indicated by Fig. 4 g and Fig. S6, CX₃CR1-deficient Gr1⁺ monocytes did not exhibit increased apoptosis rate during *Lm* infection, and the proliferation of monocytes in the spleen is not responsible for their accumulation. Total blood monocyte counts (CD115⁺ CD11b⁺ NK1.1⁻) and Gr1⁺ monocyte counts performed in Cx₃cr1⁺/⁺, Cx₃cr1⁺/-, Cx₃cl1⁺/-, and Cx₃cl1⁻/- mice did not show any difference between CX₃CR1-deficient or CX₃CL1-deficient mice and control mice (Fig. S9, a–c), and the proportion of Gr1⁺ blood monocytes that bind annexin-V in the steady state was also similar in CX₃CR1-deficient animals and in controls (Fig. S9 d). However, there was a 20% decrease in the numbers

Figure 4. CX₃CR1 is important for the recruitment of CD11b⁺ CD11c⁻ monocytes to the spleen during infection. (a and b) Mouse fractalkine/CX₃CL1 transcripts are detected in spleen tissue sections from WT mice around B cells follicles in the marginal zone/T cell area. Bars: (a) 500 μm; (b) 200 μm. (c) Short-term adoptive transfer of BM monocytes into irradiated WT recipient mice. Equal numbers (2 × 10⁵) of BM monocytes from Cx₃cr1⁺/⁺, Cx₃cr1⁺/-, Cx₃cl1⁺/-, Cx₃cl1⁺/+ mice (blue) and Cx₃cr1⁻/-, Cx₃cl1⁻/-, Cx₃cl1⁻/-, Cx₃cl1⁻/- mice (red) were mixed and adoptively transferred into a Cx₄5.1 congenic recipient. CX₃CR1-expressing CD45.1/2 Gr1⁺ monocytes are recruited in the spleen of irradiated host 10× more efficiently than CX₃CR1-deficient monocytes. The experiment was performed three times with similar results. (d–f) Monocyte recruitment in the spleen of infected mice. (d) BALB/c Cx₃cr1⁺/⁺ and Cx₃cr1⁻/- mice (seven per group) were infected i.v. with live *Lm* (3 × 10⁵), and monocytes (CX₃CR1⁺ gfp⁺ CD11b⁻ Ly-6C⁻ Ly-6G⁻, CD19⁻, CD3⁻, and NK1.1⁻) were enumerated after 24 h in the blood and spleen of infected and control uninfected mice. (e) C57BL/6 Cx₃cr1⁺/⁺ and Cx₃cr1⁻/- mice (seven per group) were injected with 7 × 10⁵ *Lm*, and monocytes were enumerated after 24 h in the blood and spleen of infected mice. (f) C57BL/6 Cx₃cr1⁺/⁺ and Cx₃cr1⁻/- mice (seven per group) were injected with 7 × 10⁵ *Lm*, and monocytes were enumerated after 48 h in the spleen of infected mice. *Lm* growth in the spleen of infected mice is shown. The asterisks indicate a significant difference between groups (P < 0.05). (g) Absence of proliferation of monocytes in the spleen of *Lm*-infected mice. BALB/c mice (n = 6 per group) were infected with 3 × 10⁵ bacteria, spleen and leg bones were harvested 16 and 48 h later, and cells were processed as indicated in Fig. S5 (available at http://www.jem.org/cgi/content/full/jem.20081385/DC1). Data indicate the percentage of BM precursors and spleen monocytes in G0/G1 and in G2+S as analyzed by flow cytometry after DNA labeling with PI. The experiment was performed three times with similar results. (h) BALB/c Cx₃cr1⁺/⁺ and Cx₃cr1⁻/- mice were infected i.v. with live *Lm* (3 × 10⁵). Data show the number of bacteria (mean ± SE) in the spleen 24 h after infection. (i) C57BL/6 Cx₃cr1⁺/⁺ and Cx₃cr1⁻/- mice were injected with 10⁴ *Lm*. Data show the number of bacteria (mean ± SE) in the spleen 24 h after infection. Circles represent individual mice.
of blood Gr1− monocytes in CX3CR1-deficient and CX3CL1-deficient mice in comparison with controls (Fig. S9, a–c), and the frequency of apoptotic Gr1− monocytes was higher in CX3CL1−/− mice in comparison with WT mice (Fig. S9 d).

The clonogenic and proliferation potentials of MDPs were then studied in vitro in the presence or absence of CX3CL1/ fractalkine. CX3CR1−/− did not confer any advantage in terms of growth or survival, even in competition with WT cells (Fig. S10, a–c, available at http://www.jem.org/cgi/content/full/jem.20081385/DC1). The role of CX3CR1 in the proliferation of MDP-derived cells in vivo was investigated by labeling MDP and CD19+ B cells, as a control, with Cell Tracker 633 (BODIPY 630/650 MeBr) and carrying out the adoptive transfer of these cells into irradiated recipients. 6 d later, 100% of both CX3CR1+/− and CX3CR1−/− MDP-derived cDCs and monocytes were no longer labeled (Fig. S10 d), indicating that they had undergone multiple rounds of division. Together with results from the adoptive transfer experiments depicted in Fig. 4, these data indicate that CX3CR1 deficiency does not significantly affect the proliferation and differentiation potential of MDP.

**DISCUSSION**

This study investigated the differentiation potential of the CX3CR1+ MDPs and the role of CX3CR1 in their development. First, our data confirmed that CX3CR1 was associated with the commitment of myeloid progenitors to the monocyte/macrophage/DC lineage. The recently described CDP (24, 25) also expresses CX3CR1, CD115, and FLT3, and its phenotype was overlapping with the MDP (this study and...
MDP can give rise to both DCs (PDCs and cDCs) and monocytes/macrophages but do not have the potential to give rise to granulocytes. Thus, they have a more restricted differentiation potential than CMPs, which can give rise to all myeloid cells, and GMPs, which can give rise to granulocytes, macrophages, and DCs (Fig. 2d). CDP has been reported to give rise to cDCs and PDCs but not macrophages and, thus, has a more restricted differentiation potential than MDPs and may be placed downstream (Fig. 2d). Further studies will be needed to investigate the developmental relationship between MDPs and CDPs.

Cx3CR1 is a later marker than CSF-1R in myeloid lineage commitment. In this study, we investigated the role of Cx3CR1/Cx3CL1 in the development of MDP-derived cells by studying mice lacking Cx3CR1 or its ligand Cx3CL1/fraction during acute and chronic inflammation and in a sensitive competitive adoptive transfer model. This study revealed an important role of Cx3CR1 in the recruitment of CD115+/Gr1+ inflammatory monocytes, which are immediate precursors to TipDCs, to the spleen during infection and for the control of bacterial growth during primary infection in two different strains of mice (C57BL/6 and BALB/c).

Impaired recruitment of Cx3CR1−deficient Gr1+ monocytes/TipDCs is the most likely mechanism underlying our observations for the following reasons: in situ hybridization experiments (Fig. 4 and Fig. S4) indicated that fractalkine is expressed in the marginal zone/T cell areas of the spleen, where TipDC accumulate during infection (Fig. 5) (44); short-term adoptive transfer of BM monocytes into irradiated hosts decreased the accumulation of Cx3CR1−deficient monocytes in the spleen by a factor of ten (see Fig. 4); during acute Lm infection, the number of monocytes was higher in the blood of infected Cx3CR1−deficient mice but decreased in the spleen (Fig. 4), and the frequency of annexin-Vhigh monocytes was not increased in spleen from infected Cx3CR1−deficient mice (Fig. S6); Cx3CR1 deficiency does not affect the survival of CD115+ Gr1+ monocytes in the periphery (Fig. S9); and finally, we have found that Cx3CR1 was redundant for MDP development, survival, homing to the spleen, and differentiation in vivo and in vitro (Fig. S10) and MDP-derived spleen cDCs and PDCs appear to develop normally.

It is of note that although CD115+ Gr1+ (Cx3CR1low) inflammatory monocytes were present in normal numbers in both Cx3CR1− and Cx3CL1−deficient mice, the present data confirm that Cx3CR1 deficiency also slightly decreases the survival of the CD115+ Gr1+ monocytes (Fig. S9). This result was expected and consistent with the conclusions that were made from our earlier adoptive transfer experiments (29) and with a more recent study (47). Cx3CR1 has been previously shown to be important for the adhesion of CD115+ Gr1− (Cx3CR1high) monocytes to the endothelium of blood vessels (38, 42). The mechanism underlying the selective survival defect of CD115+ Gr1− Cx3CR1high monocytes may thus be related to the unique patrolling behavior of these cells, which attach themselves to the luminal side of the endothelium via a Cx3CR1− and LFA1−dependent mechanism, crawling onto the endothelium for considerable periods of time without extravasation (42). Human Cx3CR1high cells also adhere to the endothelium via a Cx3CR1-dependent mechanism (38), and adhesion is known to increase survival as has been shown for Cx3CR1-expressing cells binding to Cx3CL1 via a mechanism involving an Akt/GSK-3β−mediated antiapoptotic signaling pathway (48, 49). The role of Cx3CR1 for the recruitment of CD115+ Gr1+ monocytes/TipDC precursors is reminiscent of the phenotype of CCR2−, MCP1−, and MCP3−deficient mice (45, 50). However, the disruption of the CCR2 axis results in much more dramatic phenotype because both Gr1+ monocytes exit from the BM into the bloodstream, and the recruitment to tissues of the few monocytes that exit the BM is impaired in these mice (45, 46, 50). Thus CCR2−deficient mice have reduced numbers of TipDCs. In contrast, BM output is intact in Cx3CR1−deficient mice, the number of blood Gr1+ monocytes are normal, and only their recruitment is impaired. Similarly, CCR2−deficient mice have a dramatically impaired innate response to Lm infection and die of infection within a few days (46), whereas Cx3CR1−deficient hosts exhibit a reduced clearance of Lm at 24 h after infections (Fig. 3), but at later time points we did not observe significant differences in the survival of Cx3CR1−deficient and control mice (not depicted). We therefore propose that Cx3CR1 may play an important role in mediating the recruitment of CD115+ Gr1+ inflammatory monocytes in the white pulp of the spleen during acute inflammation, possibly through adhesion to the capillary endothelial cells of the marginal zone of the spleen.

**MATERIALS AND METHODS**

**Animals.** C57BL/6 mice on C57B1/6 or C57B1/6.Cx3cr1gfp/+ congenic backgrounds were obtained from Charles River Laboratories. Cx3cr1gfp reporter mice on the C57BL/6 and BALB/c background were obtained from D. Littman’s laboratory (Skirball Institute, New York, NY). Cx3cr1−/− and C57BL/6 mice were crossed with WT C57B1/6 mice to produce Cx3cr1−/− mice on a mixed C57B1/6/C57B1/6 background for competitive adoptive transfer experiments. C57BL/6 and BALB/c Cx3cr1−/−, Cx3cr1−/− and Cx3cr1+/− and BALB/c Cx3cr1−/− and Cx3cr1gfp were bred and maintained in the specific pathogen-free animal facility of the Institut National de la Santé et de la Recherche Médicale U924. C57BL/6 and Cx3cr1gfp/+ and Cx3cr1+/+ and ApoE−/− Cx3cr1+/+ and Cx3cr1−/− mice were maintained in the laboratory of L.F. Chario (Gladstone Institute of Cardiovascular Disease, University of California, San Francisco, San Francisco, CA). All mice were maintained according to institutional guidelines and used at the age of 8–14 wk old. 4-wk-old C57BL/6 mice on the C57B1/6 background were used as adoptive transfer recipients. Experimental protocols using mice were examined and approved by the Direction Départementale des Services Vétérinaires de Paris (Paris, France).

**Antibodies and recombinant proteins.** The following purified or conjugated antibodies were purchased from BD: purified anti-FcγRIIIa/II (CD32/16, clone 2.4G2); PE-APC-, PE-P, or biotin-labeled anti-CD11b (M1/70); PE-APC-, or biotin-labeled anti-CD117 (cKit; 2B8); PE-APC-labeled anti-NK1.1 (PK136); FITC−, PE−, or APC−labeled anti-CD11c (HL3); PE−, or APC−labeled anti−TCR-B (H57-597); PE− or APC−labeled anti-B220 (RA3-6B2); biotin−labeled anti-CD45.2 (H12–2B1); PE− or APC−labeled anti-CD45.1 (H12–2B1); PE− or APC−labeled anti−TIM-3 (IMR90); FITC−, or PE−labeled antibody against CT− (38–56); PerCP− or biotin−labeled anti-CD3 (145–2C11); biotin−labeled anti-CD19 (MB19-1); APC−labeled anti−β2m (53G-1A5); PE− or APC−labeled anti−CD8−α (53–67); PerCP− or biotin−labeled anti-CD4 (RM4-5); PE− or APC−labeled anti−CD3 (145–2C11); biotin−labeled anti-CD19 (MB19-1); APC−labeled anti−TIM-3 (IMR90); or biotin−labeled anti−TIM-3 (IMR90).
APC–Alexa Fluor 750-labeled anti-CD45.2 (104), PE- or PE-Cy5-labeled anti-FcRII (A2F10), and PE- or biotin-labeled anti CD115 (AFS98). Anti-CD169 (MOMA-1) and APC- or biotin-labeled F4/80 clone CLA-1 were purchased from AbD Serotec. mPDCA1 antibody was purchased from Miltenyi Biotec. Anti-NOS-2 (M-19) polyclonal rabbit was purchased from Santa Cruz Biotechnology, Inc. Goat anti–rabbit Alexa Fluor 647 was purchased from Invitrogen. Hydroethidine was purchased from Polysciences, Inc. Antibodies used for depletion, including TER-119 (erythrocytes), RB6-8C5 (Ly6-C/G), and GK1.5 (CD4), were supernatants of hybridomas donated by B. Rocha (IFR, Necker, Paris, France). Streptavidin Pacific Blue was purchased from Invitrogen. Dead cells were gated out by PI labeling (Invitrogen). APC-labeled annexin V was purchased from BD. Recombinant mouse macrophage CSF-1 and recombinant mouse fractalkine (CX3CL1) were purchased from R&D Systems. Recombinant mouse GM-CSF was purchased from PeproTech.

Mouse blood phenotyping. Blood was drawn according to institutional guidelines, and red blood cells were lysed in 5 vol of red blood cell lysis buffer (155 mM NH4Cl, 10 mM NaHCO3, and 0.1 mM EDTA). Cell suspensions were filtered using a 40-μm filter (BD), washed twice in PBS 0.5% BSA, and an aliquot was counted with a Guava ViaCount analyzer (Guava Research). Cells were incubated with anti–mouse FcRII/III (2.4G2) for 10 min at 4°C in PBS 0.5% BSA and then stained with anti–mouse antibodies specific for CD11b (PE-Cy7), Ly6C (biotin), NK1.1 (APC), and CD115 (PE). Biotinylated antibodies were revealed by subsequent staining with streptavidin–Pacific blue. Cells were analyzed on a nine-color CYAN ADP flow cytometer (Dako) using the Summit 4.3 software (Dako), and monocytic cells were identified as CD115+, CD11b+, SSChi, NK1.1−, or Ly6c− cells. Annexin V staining was performed according to the manufacturer’s instructions. Absolute numbers were calculated using cell percentages and total white blood cells counts. Data on blood monocytes counts were obtained from littermate mice at the age of 7–8 wk for Cx3crt1+1, Cx3crt1+/−, and Cx3crt1−/− mice and 13–14 wk old mice for Cx3crt1+/+ and Cx3crt1−/− mice.

Phenotyping of splenocytes. Spleens were removed, triturated in RPMI 2% FCS at 4°C with the end of a 3-ml syringe, and then passed through a 100-μm cell strainer (BD) in one well of a 6-well plate. Cell suspensions were filtered using a 40-μm filter, washed twice in PBS 0.5% BSA, blocked with anti–mouse FcRII/III, and then stained with anti–mouse antibodies specific for CD3e, CD11c, CD19, Ly6G, CD11b, Ly6C, mPDCA1, NK1.1, and CD115. Biotinylated antibodies were revealed by subsequent staining with streptavidin–Pacific blue. Cells were analyzed on a nine-color CYAN ADP flow cytometer using Summit 4.3 software (Dako).

Phenotyping, isolation, and adoptive transfer of MDP and CDP. BM cells from femurs and tibias of Cx3crl1−/− mice aged 6 wk were flushed with serum-free RPMI. Fe receptors were blocked with antibody to FcRII/III (2.4G2), and cells were labeled using antibodies against lineage markers (IL7-Rα, CD11c, CD11b, NK1.1, CD3, Ter119, CD19, and Gr1), CD117, CD115, and CD135. For purification of MDP, red blood cells and Gr1-expressing cells were removed using Ter119 and RB68C5 antibodies as previously described (22). MDP (defined as IL7-Rα− lineage gp1 [CXCRI]) CD117+ expressing CD115; Fig. S2), CDP (defined as IL7-Rα− lineage CD135−CD115+CD117+; Fig. S3), and BM monocytes (defined as gp1− CD11b+ CD115− NK1.1− CD117−) were purified by flow cytometry using FACSVantage or FACSaria cell sorters (BD). For competitive adoptive transfer, monocytoids or MDP were purified from Cx3crt1+10 mice on a CD45.2 background and from Cx3crt1−/− mice on a CD45.2/1 background, unless otherwise indicated. FACS-purified cells from each genetic background were mixed at a 1:1 ratio and were transferred i.v. to 4-wk-old recipient mice on a CD45.1 background. Recipients were either irradiated with 10 Gy or were not irradiated. For analysis of cell populations in WT mice, Cx3crt1+10 reporter mice, and adoptive transfer recipients, blood and organs were harvested from anaesthetized mice as described previously (29), and spleens were digested with collagenase d, homogenized, and filtered. Erythrocytes were lysed using red blood cell lysis buffer for quantification of cell populations in reporter mice or were depleted by density gradient centrifugation (LSM 1077; Eurobio) for analysis of adoptive transfer recipients. PBMC or splenic mononuclear cell suspensions were enumerated with a Guava ViaCount analyzer. Fe receptors were blocked, and then cells were labeled using fluorochrome- or biotin-conjugated antibodies and analyzed by five, six, or seven-color flow cytometry using a CYAN ADP.

In vitro analysis of cell survival, proliferation, and differentiation potential. MDPs from Cx3crt1+1 or Cx3crt1−/− were purified by FACS, as described in the previous section, and plated at one cell per well into 96-well plates containing Opti-MEM (Invitrogen) supplemented with 10% FCS (PAN biotech), 100 U/ml penicillin, and 100 μg/ml streptomycin (Invitrogen) along with CSF-1, GM-CSF, or both (10 ng/ml each unless otherwise indicated) in the presence or absence of recombinant CX3CL1. Resulting colonies were enumerated after incubation at 37°C in 5% CO2 for 7 d. For differentiation analysis, colonies were recovered from wells by vigorous pipetting and were immunolabeled for cell surface markers as described in the previous section. For determination of proliferation potential, individual cells from colonies arising from single cell clones were counted using a 10 or 5× objective on a microscope (Axiowertt; Carl Zeiss, Inc.). Counting accuracy was validated by digital analysis of random colonies of various sizes photographed using a camera (CoolSnap ES; Roper Scientific) and Photoshop CS (Adobe) for marking of individual cell bodies.

For analysis of proliferation under competitive conditions, 100 MDPs each from Cx3crt1+1 or Cx3crt1−/− on different CD45. congenic backgrounds, as described in the Animals section, were deposited into wells of 96-well round-bottom plates, coated with fractalkine or BSA as control, and grown in decreasing concentrations of CSF-1. After 5 d in culture, cells were harvested from individual wells and analyzed for origin by CD45.1 and CD45.2 staining by flow cytometry. At least 10 wells were analyzed per experiment.

Lm infection. The Lm 10403s WT strain was used in these experiments. This strain exhibits an LD50 of 7 × 105 in C57BL/6 mice. Bacteria were prepared from clones grown from organs of infected mice. Stocks of bacteria were kept frozen at −80°C. For infections, bacteria were grown to a logarithmic phase (OD405 = 0.05–0.15) in Broth Heart Infusion medium (Sigma-Aldrich), diluted in PBS, and injected i.v. into lateral tail vein with 0.1–10× LD50 of WT Lm per mouse. To study production of TNF and iNOS, organs were cut in small pieces and incubated at 37°C for 20 min in HBSS medium (Invitrogen) containing 4,000 U/ml collagenase I (Invitrogen) and 0.1 mg/ml DNase I (Roche). Red blood cells were lysed for 2–3 min in 170 mM NH4Cl and 17 mM Tris HCl, pH 7.4. Cells were stained with the specified antibodies in 100 μl PBS containing 0.5% of BSA (FACS buffer). For the TNF-α intracellular staining, splenocytes were incubated at 37°C and 5% CO2 for 3–4 h in RPMI 1640 (Invitrogen) containing 5% FCS and 2 mg/ml Golgi Plug (BD) with or without 5 × 108 HKLM/ml. Cells were incubated for 20 min on ice with the indicated cell surface marker mAbs, fixed in 1% PFA (Sigma-Aldrich), dilitated in PBS, and injected i.v. into lateral tail vein with 0.1–10× LD50 of WT Lm per mouse. To study production of TNF and iNOS, cells were cut in small pieces and incubated at 37°C for 20 min in HBSS medium (Invitrogen) containing 4,000 U/ml collagenase I ( Invitrogen) and 0.1 mg/ml DNase I (Roche). Red blood cells were lysed for 2–3 min in 170 mM NH4Cl and 17 mM Tris HCl, pH 7.4. Cells were stained with the specified antibodies in 100 μl PBS containing 0.5% of BSA (FACS buffer). For the TNF-α intracellular staining, splenocytes were incubated at 37°C and 5% CO2 for 3–4 h in RPMI 1640 (Invitrogen) containing 5% FCS and 2 mg/ml Golgi Plug (BD) with or without 5 × 108 HKLM/ml. Cells were incubated for 20 min on ice with the indicated cell surface marker mAbs, fixed in 1% PFA FACS buffer for 20 min on ice, and permeabilized for 30 min in Perm/Wash (BD). For intracellular staining of TNF-α, cells were incubated for 20 min on ice in FACS buffer containing anti–TNF-α or control rat IgG. For intracellular staining of iNOS, cells were incubated for 20 min on ice in FACS buffer containing anti–iNOS rabbit polyclonal or control normal goat IgG, and staining was revealed using goat anti–rabbit Alexa Fluor 647 mAb. In all cases, cells were washed, fixed for 30 min in 1% PFA FACS buffer, and analyzed on a FACS Calibur cytosensorometer (BD). To study the production of RlO, 5 × 106 splenocytes were incubated for 3 h at 37°C and 5% CO2 in 5% FCS RPMI 1640 with 5 × 108 HKLM/ml and 160 μM hydroethidine. Hydroethidine is oxidized by RlO in red fluorescent ethidium bromide, therefore allowing for the detection
of ROI-producing cells. Cells were washed in FACS buffer and stained for expression of cell surface markers. To measure bacterial titer in the spleen, mice were injected i.v. into the lateral tail vein with 10⁴ WT Lm and organs were harvested and dissociated on metal screens in 10 ml of 0.1% Triton X-100 (Sigma-Aldrich). Serial dilutions were performed in the same buffer, and 50 μl was plated onto BHI media plates.

For cell cycle analysis, 6-wk-old BALB/c mice were infected with 3 × 10⁴ bacteria. 48 h later, spleen and leg bones were harvested. BM were flushed with RPMI 5% FCS and spleen were treated with collagenase/DNase. After lysis of red blood cells, cells suspensions were separately enriched for CD11b+ (MACs) and stained for CD11b, Ly6c, CD3, CD19, and Ly6G surface markers. Monocytes (CD11b+ Ly6c+ DX5 and stained for CD11b, Ly6c, DX5, CD3, CD19, and Ly6G surface markers.

Immunofluorescence. Spleens were fixed for 1 h in medium containing 0.05 M phosphate buffer, 0.1 M l-lysine, pH 7.4, 2 mg/ml NaIO₄, and 40 mg/ml paraformaldehyde and then dehydrated overnight at 4°C in a solution of 30% sucrose under agitation. Tissues were snap frozen in Tissue-Tek (Sakura). 10-μm frozen sections were stained with Alexa Flour 484 anti-B220 antibody (RA3-6B2; BD), Alexa Flour 647 anti-CD3 antibody (17A2; BD), and unconjugated anti-NOS2 antibody (M19; Santa Cruz Biotechnology, Inc.). NOS2 staining was revealed with an Alexa Flour 647 goat anti-rabbit IgG (Invitrogen). Immunofluorescent confocal microscopy was performed with a laser-scanning confocal microscope (TCS SP5; Leica). Final image processing was performed using ImageJ software (National Institutes of Health).

RT PCR. Standard methods were used for the preparation of total cellular RNA and first-strand complementary DNA from spleens and brains of WT C57BL/6 mice. The following primer pairs were used: Cx3cl1 forward, 5‘-CATGGGACAGATGACCTCA-3’, and reverse, 5‘-TTCCAATGTGTCGACAAGATGACCTCA-3’ (455-bp product); and cyclophilin forward, 5‘-TGGTCAACCCACCGGTGTCTC-3’, and reverse, 5‘-TCCAGCATTTGGCCTAGGACAGA-3’ (455-bp product).

In situ hybridization. The 60-mer oligonucleotide probes were synthesized and purified by Invitrogen. The oligonucleotides were 3′ end labeled with [32P]dATP (PerkinElmer) using 15 U/ml of terminal deoxynucleotidyl transferase (Invitrogen) to a specific activity of ~7 × 10⁸ cpm/μg. The probes were purified on BioSpin columns (Bio-Rad Laboratories) before use. The Cx3cl1 probes were chosen according to the human Cx3cl1 complementary DNA sequence (available under GenBank accession no. NM_009142; http://www.ncbi.nlm.nih.gov/Genbank). The sequences of the probes were 5′-CCCGAGGAGCAGGGACATACTATCTACATCTACATCATGTCTGAGCCATCTGTGATGGATTGACTTCCCTGCTATG-3′ (455-bp product); and cyclophilin forward, 5′-TGGTCAACCCACCGGTGTCTC-3’, and reverse, 5′-TCCAGCATTTGGCCTAGGACAGA-3’ (455-bp product).

The hybridization cocktail contained 50% formamide, 4× SSC, Denhardt’s solution, 0.25 mg/ml of yeast transfer RNA, 0.25 mg/ml of sheared herring sperm DNA, 0.25 mg/ml poly A+, 10% dextran sulfate (Sigma- Aldrich), 100 mmol DTT, and [32P]dATP-labeled probes (10⁵ cpm/100 μl final concentration). 100 μl of hybridization solution was applied to each section. Sections were covered with a paraffin coverslip and incubated in a humidified chamber at 43°C for 20 h. After hybridization, the slides were washed twice for 15 min in SSC supplemented with 10 mM DTT at 55°C, twice for 15 min in 0.5× SSC supplemented with 10 mM DTT at 55°C, and, finally, in 0.5× SSC supplemented with 10 mM DTT for 15 min at room temperature. The sections were dipped in water, dehydrated by incubation in a series of graded concentrations of ethanol, placed against x-ray film (Hyperfilm Betamax; GE Healthcare) for 10 d, and then against photographic emulsion (NTB2; Eastman Kodak) for 2 mo at 4°C. Sections were developed, counterstained with toluidine blue (0.2% in 0.2 M sodium acetate, pH 4.3), covered with a coverglass, and examined under bright- or dark-field illumination with a light microscope (DNRB2; Leica). Both bright- and dark-field images were collected by a charge-coupled device camera (Nikon) connected to a computer.

Online supplemental material. Fig. S1 shows the expression of CX3CR1 by PDCs. Fig. S2 shows the sorting gates for MDPs. Fig. S3 shows the effects of the AF508 anti-CD115 antibody on the response of MDP to M-CSF in vitro. Fig. S4 shows the expression of CX3CR1 in the spleen. Fig. S5 shows analysis of the proliferation of monocytes in the spleen and BM. Figs. S6 and S7 show the ratio of Gr1⁺ and Gr1⁻ monocytes in the spleens of infected and control mice and the frequency of annexin-V⁺-positive cells. Fig. S8 describes the analysis of the production of TNF, NOS2, and ROI by Gr1⁺ monocytes in the spleen of infected and controls mice. Fig. S9 indicates the absolute numbers of Gr1⁺ and Gr1⁻ monocytes in the blood of mice deficient in CX3CR1 or CX3CL1 in comparison to controls. Fig. S10 describes the role of CX3CR1 in the proliferation of MDP in vitro and in vivo.

Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20081385/DC1.

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