Hematopoietic stem cells (HSCs) reside in specialized bone marrow (BM) niches regulated by the sympathetic nervous system (SNS). Here, we have examined whether mononuclear phagocytes modulate the HSC niche. We defined three populations of BM mononuclear phagocytes that include Gr+ monocytes (MOs), Gr− MOs, and macrophages (MΦ) based on differential expression of Gr-, CD5, F4/80, and CD69. Using MO and MΦ conditional depletion models, we found that reductions in BM mononuclear phagocytes led to reduced BM CXCL2 levels, the selective down-regulation of HSC retention genes in Nestin+ niche cells, and egress of HSCs/progenitors to the bloodstream. Furthermore, specific depletion of CD169+ MΦ, which spares BM MOs, was sufficient to induce HSC/progenitor egress. MΦ depletion also enhanced mobilization induced by a CXCR4 antagonist or granulocyte colony-stimulating factor. These results highlight two antagonistic, tightly balanced pathways that regulate maintenance of HSCs/progenitors in the niche during homeostasis, in which MΦ cross talk with the Nestin+ niche cell promotes retention, and in contrast, SNS signals enhance egress. Thus, strategies that target BM MΦ hold the potential to augment stem cell yields in patients that mobilize HSCs/progenitors poorly.
Sympathetic neural tone is crucial for both steady state (Méndez-Ferrer et al., 2008) and granulocyte colony-stimulating factor (G-CSF)–enforced (Katayama et al., 2006) release of HSCs/progenitors from the BM. Recent studies indicate that mesenchymal stem cells (MSCs), identified by the expression of the intermediate filament protein Nestin, comprise a critical cellular constituent of the stem cell niche that is under the control of the sympathetic nervous system (SNS; Méndez-Ferrer et al., 2010b). Because previous studies using G-CSF receptor–deficient mice showed that expression of the receptor on transplantable hematopoietic cells was required for G-CSF–induced mobilization (Liu et al., 2000), we have previously speculated that at least two distinct pathways, neural and hematopoietic, acted in concert to promote HSC/progenitor egress (Katayama et al., 2006).

Hypothesizing that mononuclear phagocytes are crucial for stromal function of the BM, we sought to eliminate these populations to evaluate their contributions to HSC trafficking. Unexpectedly, we have found that BM macrophages (MΦ) did not promote the egress of HSCs/progenitors, but rather contributed to the retention of HSCs in the BM by acting on Nestin+MSCs. These data uncover a new role for the innate immune system in regulating stem cell niche functions.

RESULTS
Phenotypic markers of BM mononuclear phagocytes
Depletion of monocytes (MO) and/or MΦ from the BM has been accomplished with injection of clodronate liposomes (Giuliani et al., 2001) and injection of the FK-binding protein dimerizer AP20187 in transgenic Mafia mice (Burnett et al., 2004; Chang et al., 2008). Mafia mice have a Fas suicide/apoptotic system driven by the CD115 (M-CSF receptor) promoter. Previous phenotypic descriptions of BM MΦ have exclusively relied on F4/80 expression (Hume et al., 1983; Giuliani et al., 2001; Chang et al., 2008). However, this marker is also expressed on BM neutrophils (Gr-1+CD115−), Gr-1hiMO (Gr-1+CD115+), Gr-1loMO (Gr-1+CD115+; Gordon and Taylor, 2005), and eosinophils (SSCintSiglec-F−; Zhang et al., 2004; Fig. S1). To distinguish among BM mononuclear phagocytes and to elucidate their differential surface phenotypes, we purified different BM populations via cell sorting based on three markers: Gr-1 (Ly6C/G), CD115, and F4/80. As expected, neutrophil granulocytes were homogeneously represented in the Gr-1+CD115−gate (Fig. 1 A, gate I) and represented 49.6 ± 1.1% of the total BM nucleated cells. In mice, there are two subsets of CD115+MO that differen-
tially express Gr-1 (Gordon and Taylor, 2005). In concordance, the Gr-1+CD115+portion (Fig. 1 A, gate II) represented a homogenous population of MO (Fig. 1 B) that constituted 9.8 ± 0.3% of the BM and is characterized as F4/80hiCD11bhiCD68intCX3CR1intMHCIId−CD11c−CD169−(Fig. 1 C and Fig. S1 A), and will herein be termed Gr-1hiMO. The Gr-1+CD115−population (Fig. 1 A, gate III) representing 1.4 ± 0.1% of BM consisted of a population of MO (Fig. 1 B) characterized as CX3CR1hiCD11b+CD68intCD169−and will be termed Gr-1loMO.

Depletion of BM mononuclear phagocytes correlates with HSC/progenitor egress and reduction in marrow CXCL12
We first depleted BM MO/MΦ using clodronate liposome injection. 14 h after clodronate liposome administration, BM MΦ were reduced by 84% (Fig. 2 A and B), whereas Gr-1hiMO, Gr-1loMO, and total BM cellularity were reduced by 79, 88, and 24%, respectively, compared with PBS-treated animals (Fig. 2 A and C–E). Depletion of the noneosinophil Gr-1−CD115−F4/80+population with clodronate further supports the conclusion that these CD169+ cells are indeed MΦ. We assessed the effect of BM MO/MΦ depletion on circulating hematopoietic progenitors and found a marked increase in colony-forming unit activity (4.5-fold; Fig. 2 F) and Lineage–Sca-1–c-kit+ (LSK) cells (6.2-fold; Fig. 2 G) in blood after mononuclear phagocyte depletion. Notably, circulating LSK Flk2−cells, enriched in long-term repopulating HSCs (Christensen and Weissman, 2001), increased by 12.9-fold after clodronate treatment (Fig. 2 H).

We next determined the effect of mononuclear phagocyte depletion in modulating the levels of CXCL12, a chemokine released by stromal niche cells that is crucial in retention and maintenance of HSCs/progenitors in the BM (Méndez-Ferrer and Frenette, 2007). Clodronate-induced depletion of BM MO/MΦ and HSC/progenitor mobilization was associated with a 44% reduction in CXCL12 mRNA levels in total BM (Fig. 2 I) and a 40% reduction in CXCL12 protein in the BM extracellular fluid (BMEF; Fig. 2 J). Because BM MO/MΦ do not produce CXCL12 (Fig. S2), these data suggest that MO/MΦ depletion causes a reduction in CXCL12 expression by BM stromal cells.

These results were further confirmed using other conditional depletion models of mononuclear phagocytes, including transgenic mice expressing the diphtheria toxin (DT) receptor under the CD11b promoter (CD11b-DTR;
47% reduction in BMEF CXCL12 (Fig. S3 S). These data are consistent with a recent study also using liposomal clodronate and Mafia mice that demonstrated the association of BM mononuclear phagocyte depletion with HSC/progenitor egress (Winkler et al., 2010). Progenitor release cannot be explained by nonspecific cell death because in vivo depletion of neutrophils and Gr-1hi MO using anti–Gr-1 (Ly6G/C) antibody or depletion of dendritic cells using DT administration in CD11c-DTR mice (Jung et al., 2002) did not lead to any progenitor mobilization (unpublished data). Collectively, these results suggest that mononuclear phagocytes play a critical role in the retention of HSCs/progenitors in the BM.

Mononuclear phagocytes regulate Nestin+ niche cells that maintain HSCs in the BM

Recent studies have revealed that GFP expression, when driven by Nestin regulatory elements, identifies rare MSCs that form HSC niches (Méndez-Ferrer et al., 2010b). We observed CD68+ and CD169+ cells throughout the BM and in the vicinity of rare Nestin+ MSC niche cells (Fig. 3 A). To evaluate whether mononuclear phagocytes regulate Nestin+ niche cells, we used clodronate liposomes to deplete BM MΦs (Fig. 1A). Treatment of CD11b-DTR chimeric animals with DT reduced MΦ counts by >40% (Fig. S3 A) and MO subsets by >50% (Fig. S3, B and C). Consistent with the expression of CD11b, also on neutrophils, total BM cellularity was reduced by ~40% in this model (Fig. S3 D). Concomitantly, progenitors circulating in blood were significantly increased by ~1.6-fold (Fig. S3 E) and BMEF CXCL12 was significantly reduced (Fig. S3 F). AP20187-treated Mafia BM chimeras depleted BM MΦ, MO, and total BM cells by 40, >80, and 30%, respectively (Fig. S3, G–J). This was associated with an ~6-fold increase in circulating progenitors (Fig. S3 K), and 46% reduction in BM CXCL12 levels (Fig. S3 L). In nontransplanted Mafia mice, mobilization was even more robust; AP20187-treated Mafia animals exhibited a >80% reduction in MO/MΦ (Fig. S3, M–P), 31% reduction in BM cellularity (Fig. S3 Q), >20-fold increase in circulating progenitors (Fig. S3R), and...
controls (Fig. S4, D–G). These results, together with the gene expression analyses (Fig. 3, B–E; and Fig. S5), suggest that BM mononuclear phagocytes play a role in HSC/progenitor retention by regulating maintenance of retention gene expression specifically in Nestin+ niche cells, but not osteoblasts.

Microarray expression analyses have shown that Nestin+ cells express high levels of Csf1 (Méndez-Ferrer et al., 2010b), a critical cytokine for MΦ development and survival (Hamilton, 2008). To determine whether Nestin+ cells in the BM regulate mononuclear phagocyte numbers, we depleted Nestin+ cells by administering DT into tamoxifen-treated Nes-CreERT2/iDTR animals (Méndez-Ferrer et al., 2010b). We found no difference in BM MΦ (Fig. S6 A) or MO (Fig. S6, B and C) in Nes-CreERT2/iDTR mice treated with tamoxifen and DT, compared with control iDTR animals.

BM MΦ produce a protein factor that raises CXCL12 production by stromal cells in vitro

The aforementioned results demonstrate a robust in vivo correlation between BM MO/MΦ depletion, and HSC/progenitor mobilization. To further dissect the effect of mononuclear phagocytes on stromal cell function, we established long-term murine Dexter BM cultures consisting of an adherent stromal layer and attached hematopoietic cells (Dexter et al., 1977). Consistent with the in vivo data, we found that clodronate liposome treatment of Dexter cultures dramatically reduced the cellularity in the wells, especially among the adherent MΦ, compared with PBS liposome-treated wells (Fig. 4 A). The number of CD115+ mononuclear phagocytes was indeed reduced by >50% after liposomal clodronate treatment (Fig. 4 B). These changes in mononuclear phagocytes were associated with decreased stromal production of CXCL12 at 24 h (~30%) and 72 h (~40%); Fig. 4 C).
Because adherent MΦ appear to interdigitate the stroma in Dexter cultures, we sought to determine the relative contribution of MΦ and MO in stromal cell–mediated progenitor retention. Thus, we used the murine stromal cell line MS-5 to determine whether the addition of MΦ or MΦ-synthesized products could affect the stromal niche. MS-5 stromal cells have been used as an appropriate in vitro model in which to replicate in vivo modulation of CXCL12 production by MΦ in the BM. When grown in medium conditioned by the MΦ cell line RAW264.7 (Raschke et al., 1978), CXCL12 production by MS-5 was significantly increased (Fig. S7A), whereas no significant difference was observed in medium conditioned by the myeloid myeloblast cell line M1 (Ralph et al., 1983; Fig. S7B). Co-culture of BM-derived MΦ (BMDM) or media conditioned by BMDM increased MS-5 production of CXCL12, indicating that a secreted soluble factor induces CXCL12 up-regulation by MS-5 cells (Fig. 4, D and E). This secreted factor was a protein, as digestion with Proteinase K abrogated the ability of MΦ-conditioned medium to raise CXCL12 production by MΦ (Fig. 4 F). We have evaluated putative candidate factors using antibodies or knockout animals for IGF-1, IL-1, TNF, or IL-10. However, the loss-of-function of any one of these factors did not alter the ability of BMDM-conditioned medium to induce CXCL12 synthesis (Fig. S7, C–F). These data suggest that BM MΦ, through the secretion of a yet undefined protein factors, directly promote the retention of HSCs/progenitors by raising CXCL12 production in BM niche cells.

Depletion of BM CD169+ MΦ mobilizes HSCs/progenitors

The MS-5/MΦ co-culture experiments suggest that differentiated BM MΦ, rather than MO, are the mononuclear phagocytes that promote HSC/progenitor retention. To test directly whether BM MΦ are promoting HSC/progenitor retention in the BM, we took advantage of the differential expression of CD169 between MO and MΦ and mice expressing DTR under the endogenous CD169 promoter (Miyake et al., 2007). Treatment of heterozygous CD169-DTR mice with DT-depleted MΦ (Fig. 5, A and B), but not MO (Fig. 5, A, C, and D), in the BM. DT treatment was associated with a 3.5-fold increase in circulating hematopoietic progenitors, as assessed by LSK cell enumeration (Fig. 5 E), and a 5.4-fold increase in the stem cell-enriched LSKFlk2+ fraction (Fig. 5 F). Moreover, depletion of CD169+ MΦ from long-term Dexter culture resulted in a 42% reduction in the ability of BM stromal cells to produce CXCL12 (Fig. 5 G). Thus, CD169+ MΦ in the BM promote stromal production of CXCL12, and their specific depletion in vivo is sufficient to mobilize HSCs/progenitors.

Parallel and antagonistic roles of MΦ and SNS in regulating HSC/progenitor release

The SNS is crucial in HSC/progenitor trafficking (Katayama et al., 2006; Méndez-Ferrer et al., 2008) where β3-adrenergic receptor (β3R) signaling plays a key role in circadian oscillations of HSC release, and both β2-adrenergic receptor (β2R) and β3R signaling cooperate in G-CSF–enforced egress (Méndez-Ferrer et al., 2010a). MΦ could act independently, or alternatively, through alterations in the sympathetic tone. Thus, to assess whether MΦ operate through a distinct pathway, we examined whether clodronate treatment was capable of inducing mobilization in sympathectomized animals. We found that mice chemically sympathectomized with 6-hydroxydopamine (6OHDA) still exhibited significant progenitor mobilization (>7-fold) in response to clodronate.
treatment; however, the absolute number of mobilized progenitors did not reach the level of SNS-intact Clodronate-treated mice (Fig. 6 A, mid-left bars). To evaluate this issue using another model, we used mice deficient in $\beta2R$ ($Adb2^{-/-}$) treated with an antagonist to the $\beta3R$. Clodronate treatment was still able to cause HSC/progenitor mobilization (>3-fold) in these mice, but again HSCs/progenitors did not mobilize to the same level as wild type animals (Fig. 6 A, mid-right bars). To dissect further the relative contribution of the $\beta2R$ and $\beta3R$ in SNS promotion of HSC/progenitor release, we evaluated the effect of $\text{M}\Phi$ depletion in mice singly lacking $\beta2R$. (Fig. 6 A, right bars) or $\beta3R$. (Fig. 6 B). Whereas clodronate treatment led to a robust increase in circulating progenitors in $Adb2^{-/-}$ animals compared with wild-type animals (Fig. 6 A, right bars), the response was blunted in $Adb3^{-/-}$ mice (Fig. 6 B). These data are consistent with previous studies demonstrating that $\beta3R$, but not $\beta2R$, signaling is critical for physiological HSC/progenitor release (Méndez-Ferrer et al., 2008). The fact that $\text{M}\Phi$ depletion can still mobilize HSCs—albeit at lower amplitude—when SNS signaling is disrupted, argues that the SNS and the BM $\text{M}\Phi$ act through distinct parallel pathways. Thus, these data suggest antagonistic functions of the autonomic nervous system and innate immunity in regulating the niche (Fig. 6 C), where the SNS promotes egress by reducing the expression of key retention factors by the niche cell (Méndez-Ferrer et al., 2010b), and in contrast, BM $\text{M}\Phi$ promotes the expression of these genes and HSC/progenitor retention in the BM.

**BM $\text{M}\Phi$ depletion synergizes with enforced HSC/progenitor mobilization**

Because the expression of retention genes is still reduced in the Nestin$^+$ niche cells 7 d after clodronate treatment (Fig. S5), we sought to determine the duration of $\text{M}\Phi$ reduction and the kinetics of recovery. We found that BM $\text{M}\Phi$ remain markedly (>90%) reduced 10 d after clodronate treatment (Fig. 7 A). Recovery started by day 16 (58% reduction) and clodronate-treated mice demonstrated no reduction in $\text{M}\Phi$ counts by day 28. Alternatively, MO populations in the BM began their recovery by day 7 after clodronate administration (unpublished data).

Interestingly, levels of LSK (Fig. 7 B) and LSKFlk2$^-$ (Fig. 7 C) inversely matched BM $\text{M}\Phi$ counts. HSCs/progenitors were still elevated in the circulation 10 d after clodronate administration (4.2- and 3.2-fold, respectively), and mobilization persisted at least until day 16 (Fig. 7 C). These data further support the specific role of BM $\text{M}\Phi$, but not MO, in promoting the retention of HSCs/progenitors in the BM.

Because BM $\text{M}\Phi$ promote HSC/progenitor retention, we examined whether elimination of this population would enhance mobilization using the CXCR4 antagonist AMD3100 or G-CSF, both of which are clinically approved mobilizing agents. We found that clodronate treatment 14 h before harvest doubled HSC/progenitor mobilization in AMD3100-treated animals and resulted in a significant increase in the number of HSCs/progenitors mobilized by G-CSF (Fig. 7, D–E). To further assess the role of $\text{M}\Phi$ in a situation where progenitor mobilization is suboptimal, we treated mice with G-CSF (Hidalgo et al., 2004) for 2 d, instead of 4 d. We found that CFU-C, LSK, and LSKFlk2$^-$ cells in the peripheral blood mobilized by 2 d of G-CSF were increased 3.8-, 5.9-, and 9.3-fold (Fig. 7, F–H), respectively, in mice that were treated with clodronate liposomes 14 h before harvest. Furthermore, we tested whether depleting $\text{M}\Phi$ 10 d before blood collection, rather than 14 h before collection, could synergize with 4 d G-CSF treatment. Indeed, when mice were preinfused with clodronate liposomes 10 d before harvest, they had 5.2-, 3.4-, 2.7- fold higher CFU-C, LSK, and LSKFlk2$^-$ in the peripheral blood, respectively (Fig. 7, I, J, K). Thus, targeting BM $\text{M}\Phi$ may be a novel modality by which to enhance mobilization yields in patients.

**DISCUSSION**

In this study, we sought to identify the role of BM mononuclear phagocytes in HSC/progenitor mobilization. Unexpectedly, we found that depletion of mononuclear phagocytes...
of niche cells that express Nestin (Méndez-Ferrer et al., 2010b).

Although osteoblasts have been proposed to represent a HSC niche, selective modulation of osteoblast numbers do not necessarily alter HSC numbers (Wilson and Trumpp, 2006; Kiel et al., 2007; Zhu et al., 2007), and the lack of osteoblasts in sites of extramedullary hematopoiesis suggest that they are dispensable to support HSCs. Recent studies have suggested that a more primitive precursor of osteoblasts compose the stem cell niche (Méndez-Ferrer et al., 2010b; Omatsu et al., 2010). Steady-state Nestin+ MSCs express substantially higher levels of genes required in HSC/progenitor maintenance and retention, including Cxcl12, compared with in vivo sorted (Fig. 3 and Fig. S5) or cultured osteoblasts (Méndez-Ferrer et al., 2010b). The relatively low expression of Cxcl12 detected in sorted osteoblasts is consistent with a recent study showing that DT administration into Cxcl12-DTR-GFP mice did not result in loss of spindle-shaped N-cadherin–expressing osteoblasts (Omatsu et al., 2010). It has been proposed that MΦ depletion mobilizes HSCs/progenitors by disruption of the

using four in vivo models was sufficient to mobilize HSCs/progenitors. Before this study, mononuclear phagocytes in the BM had been poorly characterized and relied on F4/80, a marker with a promiscuous expression profile in the BM (Fig. S1). Herein, we rigorously discriminated among Gr−1hi MOs, Gr1lo MOs, and MΦ in the BM by differential expression of Gr−1, CD115, F4/80, CD11b, CD11c, MHC class II, CX3CR1, CD68, and CD169. Using CD169DTR/+ animals, we were able to implicate MΦ, constituting ~2.6% of total BM cells, in the promotion of HSC/progenitor retention through interaction with the recently characterized population

Figure 5. Depletion of BM CD169+ MΦ, but not CD169− MOs, mobilizes HSCs/progenitors.

(A–D) Wild-type (WT) or heterozygous CD169-DTR (CD169DTR/) mice were treated with DT. (A) Representative dot plots show the percentages of BM mononuclear phagocytes in wild-type (top) or CD169DTR/+ mice treated with DT (bottom). (B–D) Bar graphs depict the absolute numbers of mononuclear phagocytes (n = 6) in wild type mice and wild type or CD169DTR/+ mice injected with DT. (E–F) Bar graphs enumerate Lineage− Sea-1− c-kit− (LSK; E) and LSKFlk2− cells (F) per milliliter of blood in the same mice analyzed in B–D (n = 6). Data are pooled from two independent experiments. (G) Cxcl12 levels were measured 72 h after administration of PBS or 1 µg/ml DT into Dexter cultures plated from the BM of CD169-DTR animals (n = 3–4 wells). Representative data from two independent experiments are shown.

Figure 6. Opposite influences of the MΦ and the SNS on HSC/progenitor retention.
(A) CFU-C after treatment with PBS (blue bars) or clodronate liposomes (red bars) in wild-type C57BL/6 (n = 11–13), 6OHDA-treated mice (n = 12–13), β2-adrenergic receptor–deficient (Adrb2−/−) mice treated with an antagonist to the β3-adrenergic receptor (Adrb3R; n = 10), and Adrb2−/− mice (n = 7). Data are pooled from three independent experiments and analyzed with one-way ANOVA/Neuman-Keuls test. (B) CFU-C after treatment with PBS (blue bars) or clodronate (red bars) in wild-type FVB mice (n = 5) and β3-adrenergic receptor-deficient (Adrb3−/−) mice (n = 8). Data are pooled from two independent experiments. (C) Schematic of antagonistic regulation of HSC/progenitor retention by MΦ and the SNS.
Figure 7. MФ depletion synergizes with AMD3100 and G-CSF mobilization. (A–C) Kinetics of MФ reduction (A) and LSK (B) and LSKFlk2− (C) mobilization, at the indicated time points, after administration of PBS- (blue) or clodronate-encapsulated (red) liposomes (n = 3–4). Data are pooled from two independent experiments. (D–E) CFU-C from peripheral blood of mice that were treated with PBS (blue) or clodronate liposomes (red; 14 h before harvest) and mobilized with AMD3100 (D; 1 h before harvest) or G-CSF (E; 4 d). Data are pooled from two independent experiments. (F–H) CFU-C (f), LSK (g), and LSKFlk2− (h) cells from the peripheral blood of mice that were mobilized for 2 d with G-CSF and treated with PBS- or clodronate-encapsulated liposomes 14 h before harvest. Experiment was performed once (n = 4), (I–K) CFU-C (I), LSK (J), and LSKFlk2− (K) from 16-wk-old female mice that were pretreated with PBS- or clodronate-encapsulated liposomes 10 d before harvest and mobilized with G-CSF for 4 d. Data are representative of two independent experiments (n = 4).

osteoblasts either 14 h or 7 d after the depletion of MФ, despite the fact that HSCs/progenitors were clearly elevated at both time points. In contrast, significant reductions in the expression of HSC retention genes were observed in Nestin+ cells, and correlated with persistent HSC/progenitor mobilization even 7 d after clodronate treatment. These results thus indicate that CD169+ BM MФ promote HSC retention by acting specifically on the Nestin+ HSC niche in the BM.

Previous studies have revealed that a transplantable cell expressing the G-CSF receptor (G-CSFR) is essential for G-CSF-induced mobilization (Liu et al., 2000). An accompanying study, using an elegant mouse model in which the G-CSFR is expressed exclusively in CD68-expressing cells, also implicates mononuclear phagocytes in G-CSF mobilization (see Christopher et al. in this issue). Thus, G-CSF signaling exclusively in MФ is sufficient to reduce niche retention and promote HSC/progenitor mobilization. However, because G-CSF-induced HSC/progenitor mobilization is at least three times more potent than MФ depletion with clodronate liposomes (this study; Winkler et al., 2010), it must also be acting on cells other than MФ.

The SNS is required for progenitor egress (Katayama et al., 2006; Méndez-Ferrer et al., 2010a), suggesting that G-CSF–mediated increase of sympathetic tone in the BM may represent a putative MФ-independent target. Our results using models with impaired sympathetic activity suggest that BM MФ exert antagonistic, independent regulatory functions in the HSC niche compared with the SNS. Although G-CSF likely has several targets in the BM microenvironment, the present data uncover two distinct opposing activities that lead to major changes in HSC retention by Nestin+ niche cells. We thus propose that G-CSF induces HSC mobilization by inhibiting MФ-mediated retention signals and simultaneously enhancing sympathetic-mediated progenitor release (Fig. 6 C).

These results expand our understanding of HSC niche components by implicating a cellular constituent of the innate immune system, the CD169+ MФ, as a niche regulator. Because targeted reduction of BM MФ can enhance HSC/progenitor mobilization, the use of antibodies against CSF-1 (M-CSF), its receptor, or small molecule inhibitors of M-CSF signaling may provide a novel strategy to increase the efficiency of HSC/progenitor mobilization for autologous transplantation.

MATERIALS AND METHODS

Mice. All experiments, unless otherwise noted, were performed on 8–10-wk-old C57BL/6 male mice from Charles River Laboratories (Frederick Cancer Research Center, Frederick, Maryland). B2-adrenergic receptor-deficient (Adhr2tm1Hof/J; Chruscinski et al., 1999; gift from G. Karsenty, Columbia University, New York, NY), B3-adrenergic receptor-deficient (FVB/N-Adhr3tm1Cof/J; Susulic et al., 1995; The Jackson Laboratory), FVB/N-CD11b-DTR (Cailhier et al., 2005; gift from C. Alomar, Mount Sinai Medical Center, New York, NY), Mφ (C57BL/6-Tg[Csf1r-EGFP-NGFR/FKBP1A/TNFRSF6][2Bck/J; Burnett et al., 2004; The Jackson Laboratory], CX3CR1/GFP [Jung et al., 2000; gift from D. Littman, New York University, New York, NY], CD169-DTR (Miyake et al., 2007), Nes-Cnp (Mignone et al., 2004), Nes-CreER2 (Balordi and Fishell, 2007), iDTR (C57BL/6-Tg[Rosa26Sortm1[HBEGF]Awaj/J; Yagi et al., 2005), IL-10−/− (Berg et al., 1996;
gift from H. Xiong, Mount Sinai Medical Center, New York, NY), and TNF-α (Marno et al., 1997; gift from M. van den Brink, Memorial Sloan Kettering Cancer Center, New York, NY) mice were also used in these studies. FVB/N, CD11b-DTR, and Mafia (CD45.2) BM chimeras were generated by transplanting 2.0 × 10⁶ and 1.9 × 10⁶ BM-nucleated cells from male donors into lethally irradiated 8-wk-old FVB/N (The Jackson Laboratory) and C57BL/6 Ly5.2 (CD45.1) male mice, respectively. BM and blood of Mafia BM chimeras showed >95% donor chimerism as assessed by flow cytometry 1 mo after transplantation. Mice were maintained on a 12 h light/12 h darkness lighting schedule. All in vivo experiments were harvested between 12:00 and 1:00 p.m. (Zeitgeber time 5:00 and 6:00) to limit circadian variations in HSC/progenitor release (Méndez-Ferrer et al., 2008) and mobilization (Lucas et al., 2008). All mice were housed in specific pathogen-free facilities at the Mount Sinai School of Medicine or Albert Einstein College of Medicine animal facility. Experimental procedures performed on the mice were approved by the Animal Care and Use Committee of the Mount Sinai School of Medicine and Albert Einstein College of Medicine.

In vivo cell depletion. C2M NDP (or clodronate) was a gift from Roche (Van Rooijen and Sanders, 1994). Clodronate liposomes (250 μl) were infused i.v. at 1 d (14 h), 7 d, 10 d, 16 d, or 28 d before harvest. CD11b-DTR mice were transplanted with DT i.p. 25 ng/kg 8 h before harvest. DT was purchased from Sigma-Aldrich. AP20187 was a gift from Ariad Pharmaceuticals. Lyophilized AP20187 was dissolved in 100% ethanol at a concentration of 62.5 mg/ml stock solution and was stored at –20°C. As recommended by Ariad Pharmaceuticals, injection solutions were prepared with a diluent composed of 4% ethanol, 10% PEG-400, and 2% Tween-20 in water. All injections were administered i.v. within 30 min after preparation. The volume of injection solution was adjusted according to the average mouse body weight to deliver a dose of 10 mg/kg AP20187 per mouse in an mean volume of 100 μl. Mice were injected daily for 5 d before harvest. Heterozygous CD169-DTR (CD169(DTR–/–)) and control C57BL/6 were injected i.p. with 10 μg/kg DT 48 h before harvest. Depletion of Nestin+ cells was accomplished as previously described (Méndez-Ferrer et al., 2010b).

Flow cytometry and cell sorting. Fluorochrome-conjugated or biotinylated mAbs specific to mouse Gr-1 (Ly6G/C, clone RB6-8C5), CD115 (clone AF598), Siglec-F (clone E50-2440), CD11b (clone M1/70), CD11c (clone N418), I-A/E (clone M5/114.15.2), CD45 (clone 30–F11), Sca-1 (clone D7), Flk2 (clone A2F10), CD117 (clone 2B8), CD3 (clone 14-5C11), B220 (clone RA3-6B2), Ter119 (clone TER-119), CD51 (clone RMV-7), and CD31 (clone MECA-13.1), corresponding isotype controls, and secondary reagents (efluor450-, APC–efluor780–, and PE–Cy7–conjugated streptavidin) were purchased from eBioscience. Anti-F4/80 (clone C11/186.1), CD68 (clone FA-11), and CD169 (clone 3D6.112) were purchased from AbD Serotec. CD68 was stained extracellularly and subsequently intracellularly with the Cytofix/Cytoperm kit (BD) according to the manufacturer’s protocol. Multiparameter analyses of stained cell suspensions were performed on an LSR II (BD) and analyzed with FlowJo software (Tree Star, Inc.). To isolate Nestin+ and Nestin– cells from the BM for Q-PCR, RBC-lysed BM cells were digested with collagenase, trypsin, and DNase, as previously described (Méndez-Ferrer et al., 2010b). Endothelial cells and osteoblasts were isolated similar to previous studies (Semerad et al., 2005; Winkler et al., 2010). In brief, tibias, femurs, and humeri of mice were flushed thoroughly of BM cells, chopped with a scalpel, and washed three times through a 5-ml polystyrene tube with blue-top cell strainer (BD) to further remove residual BM cells. The bone fragments were then digested at 37°C with Type IA collagenase (Sigma–Aldrich) for 40 min while spinning. RBC-lysed pellet was then stained for sorting. Cells were sorted by Moflo Cell sorter (Dako) at the Flow Cytometry Core Facility at Mount Sinai School of Medicine or Aria Cell sorter (BD) at the Flow Cytometry Core Facility at Albert Einstein College of Medicine.

CFU-c assays and mobilization. Colony-forming assays were performed as previously described (Frenette et al., 1998). Mobilization experiments with AMD3100 and 4-d G-CSF were performed as previously described (Katayama et al., 2006; Lucas et al., 2008). Some experiments were performed with suboptimal G-CSF treatment (2 d), as previously described (Hidalgo et al., 2004).

RNA isolation, reverse transcription, and Q-PCR. For measurement of BM Cxcl12 gene expression, femurs were flushed and mixed with 0.5 ml Trizol (Invitrogen) and stored at −80°C. Conventional reverse transcription, using the Sprint PowerScript reverse transcription (Takara Bio Inc.) was performed in accordance with the manufacturer’s instructions. Q-PCR was performed with SYBR GREEN on an ABI PRISM 7900HT Sequence Detection System (Applied Biosytems). The PCR protocol consisted of one cycle at 95°C (10 min) followed by 40 cycles of 95°C (15 s) and 60°C (1 min). Expression of Gapdh was used as a standard. The mean threshold cycle number (Cq) for each tested mRNA was used to quantify the relative expression of each gene: 2^(-Cq[Gapdh] - Cq[gene]). Primers used are listed below: Cxcl12_FWD, 5′-CGCAAGGTCGTGCACGCG-3′; Cxcl12_REV, 5′-TTGGCCTGCGGATGTTGCG-3′; Ang1_FWD, 5′-CTGCTCAGCACCTCATCATCA-3′; Ang1_REV, 5′-CACTTCTTATTTGACCGAGCT-3′; Kitl_FWD, 5′-CCCTGAAGACCTGGCCTA-3′; Kitl_REV, 5′-CAATTCAAGGAAATGAGGACC-3′; Vcam1_FWD, 5′-GAGCTGTTCGACGGGGTCTA-3′; Vcam1_REV, 5′-CTTCCATCCTATAGCAATAGGTTG-3′; Osteocalcin_FWD, 5′-GGCGCAATAAGCTGTTGACAG-3′; Osteocalcin_REV, 5′-GGCAGCACAGGCTCTAAAGTT-3′; Gapdh_FWD, 5′-TGTTGTCCTGCTGGATCTGTA-3′; Gapdh_REV, 5′-CCCTGCTTACACCATCCTTTCA-3′.

CXCL12 ELISA. 96-well ELISA plates were coated overnight at 4°C with 50 μl of 2 μg/ml anti-CXCL12 coating antibody (MAB530; R&D Systems). Next, the wells were washed three times with wash buffer (0.05% Tween 20 in PBS) and incubated for 1 h at room temperature with 200 μl of blocking buffer (1% BSA, 5% D-Sucrose, and 0.05% NaCl, in PBS, all from Thermo Fisher Scientific). After 5 washes, 100 μl of samples diluted 1:2 in PBS were added and incubated for 2 h at room temperature. After 3 washes, 100 μl of 0.250 μg/ml polyclonal biotinylated anti-human/mouse SDF-1 (BAF310; R&D Systems) was added and incubated for 30 min. After 3 additional washes, the reaction was developed by incubation for 30–30 min with 50 μl of TMB substrate solution (Sigma–Aldrich) and stopped by adding 50 μl of 1M HCl solution (Thermo Fisher Scientific). Optical density was determined with a microplate reader set at 450 nm. Optical density of PBS control wells was subtracted from optical density of samples. Reconstituant mSDF-1x (PeproTech) was used to generate a linear standard curve.

Immunofluorescence. Anesthetized Nε–GFP transgenic animals (Mignone et al., 2004) were perfused, and femurs and tibia were sectioned and stored as previously described (Méndez–Ferrer et al., 2008). Slides were washed three times in Coplin jars (Sigma–Aldrich) to remove OCT solution residue, and then incubated for 1 h at room temperature in 20% goat serum (Sigma–Aldrich) diluted in PBS + 0.1% Tween solution (PBS-Tw). After 3 washes in PBS-Tw, slides were incubated in PBS-Tw + 2% goat serum + 0.5% Triton X-100 for 1 h at room temperature. After 3 washes in PBS-Tw, slides were incubated in primary antibody (anti-CD68–Alexa Fluor 647 [clone FA-11; AbD Serotec] and anti–CD169–biotin [clone MOMA-1; AbCam]) at 1:1000 concentration in PBS-Tw + 2% goat serum overnight in the dark at room temperature. After three washes in PBS-Tw, CD169 staining was continued with streptavidin–PE staining for 5 min. Slides were mounted with Vectashield + DAPI, covered, and sealed with nail polish. Images were acquired on an Examiner microscope (Carl Zeiss, Inc.) and all images were processed using Slidebook software (Intelligent Imaging Innovations, Inc.).

Long-term BM Dexter cultures. 3.7 × 10⁶ BM-nucleated cells were plated in 1 ml Dexter medium [Myelocult M5300 media [Stem Cell Technologies]] supplemented with 1% penicillin–streptomycin [Cellgro],
1% amphotericin [Cellgro], and 10⁻⁵ M freshly thawed Hydrocortisone [Sigma-Aldrich] in 12-well plates. Cultures were maintained in a water-jacketed incubator at 33°C and 5% CO₂. Half the media was changed weekly for 6 wk.

In the sixth week, the culture media was removed and replaced with 1 ml of 3% DEXTRG media containing 40% PBS- or chlordane-encapsulated liposomess by volume in some experiments. After 24-h incubation, the culture media was removed and 1 ml fresh media was added. 24 h and 72 h later, the media was collected and frozen to assess CXCL12 levels by ELISA and the adherent layer was Hema 3-stained to assess cell morphology or detached by cell scraper (BD) for flow cytometric analysis. In experiments with DEXTRG cultures derived from CD169DTR/+ BM, the media was removed and replaced with 1 ml fresh DEXTRG medium containing PBS or 1 µg/ml DT. After 72 h of incubation, the media was frozen and later assessed for CXCL12 levels by ELISA.

MS-5 cell culture. MS-5 cells were grown in monolayers in complete medium (α-MEM medium supplemented with 10% FBS [Stem Cell Technologies], penicillin-streptomycin [Invitrogen], 5% glucose [Invitrogen], and 5% sodium pyruvate [Invitrogen]). Cultures were maintained at 37°C and 1:10 split with 0.05% trypsin-EDTA (Invitrogen) every 3 or 4 d, when cells reached ~80% confluence: 5,000 MS-5 cells were plated in 300 µl complete medium in 48-well plates for 24 h before addition of BMDM or medium conditioned by BMDM (MФ CM), RAW264.7, or M1 cells (see below).

BM MØ/MФ cell culture. The M1 myeloblast cell line was cultured in DME medium (Cellgro) supplemented with 10% FBS, 2 mM l-glutamine (Sigma-Aldrich), 1% sodium pyruvate (Invitrogen), and 1% penicillin-streptomycin. RAW264.7 MФ cell line (a gift from B. Tenover, Mount Sinai Medical Center, New York, NY) was cultured identically to the M1 myeloblast cell line, except for addition of 100 mM Hepes (Sigma-Aldrich). 50,000 RAW264.7 and M1 cells were plated in 300 µl in a 48-well plate. 3 d later, the supernatant was collected and centrifuged at 15,000 g to remove cellular debris, mixed with MS-5 complete media at 25 or 50% concentration and added to MS-5 cells (see previous paragraph). BMDM were derived by plating 10,000 total BM-nucleated cells in 300 µl of complete medium with 25% conditioned medium from the sarcoma cell line MS-180 (Rosenthal et al., 1990). The MS-180–conditioned medium was observed to generate MФ CM by flow cytometry. In some experiments, BMDMs were derived from MS-5 complete media (see previous paragraph).

Proteinase K digestion of MФ CM. MФ CM was treated with 0.5 mg/ml proteinase K (Sigma-Aldrich, St. Louis, MO) for 30 min at 37°C and then heat inactivated at 95°C for 10 min. Heat-inactivated FBS was re-added to MФ CM to the 10% concentration of the native media.

Sympathectomy. Chemical sympathectomy was performed by i.p. injection of 100 mg/kg and 250 mg/kg of 6OHDA (Sigma-Aldrich) in PBS solution containing 200 µg/kg and 500 µg/kg acetic acid (Sigma-Aldrich) on day -4 and -2, respectively, before harvest. Pan-adrenergic receptor abrogation was accomplished by injecting Adiy2m165/J (Chruscinski et al., 1999) animals with 2.5–5 mg/kg of β3-adrenergic receptor antagonist (SR59230A, i.p.; Sigma-Aldrich).

Statistical analyses. Unless otherwise indicated, the unpaired Student’s t test was used in all analyses, data in bar graphs are represented as mean ± SEM, and statistical significance was expressed as follows: * P < 0.05, ** P < 0.01; *** P < 0.001; ns, not significant

Online supplemental material. Fig S1 shows that BM neutrophils, MOs, and eosinophils express F4/80. Fig S2 shows that MФ do not produce CXCL12. Fig S3 shows three other models in which mononuclear phagocyte depletion is associated with robust HSC/progenitor mobilization and CXCL12 reduction. Fig S4 shows the sorting strategy for Nestin” and Nestin” fractions and bone endothelial cell and osteoblast fractions. Fig S5 shows that retention gene expression is reduced in Nestin” cells seven days after mononuclear phagocyte treatment. Fig S6 shows that mononuclear phagocytes are not reduced 7 d after depletion of Nestin” cells. Fig S7 shows that soluble factor from a MФ, but not myeloblast, cell line enhances stromal CXCL12 production. Online supplemental material is available at thompson-snipes, m.w., leach, d.m., and rennick, m. 1996. enterocolitis in mice: a model for human inflammatory bowel disease. Proc. natl. acad. sci. u.s.a. 93:15525–15529. doi:10.1073/pnas.931552593


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