Macrophages retain hematopoietic stem cells in the spleen via VCAM-1

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Splenic myelopoiesis provides a steady flow of leukocytes to inflamed tissues, and leukocytosis correlates with cardiovascular mortality. Yet regulation of hematopoietic stem cell (HSC) activity in the spleen is incompletely understood. Here, we show that red pulp vascular cell adhesion molecule 1 (VCAM-1)+ macrophages are essential to extramedullary myelopoiesis because these macrophages use the adhesion molecule VCAM-1 to retain HSCs in the spleen. Nanoparticle-enabled in vivo RNAi silencing of the receptor for macrophage colony stimulation factor (M-CSFR) blocked splenic macrophage maturation, reduced splenic VCAM-1 expression and compromised splenic HSC retention. Both, depleting macrophages in CD169 iDTR mice or silencing VCAM-1 in macrophages released HSCs from the spleen. When we silenced either VCAM-1 or M-CSFR in mice with myocardial infarction or in ApoE−/− mice with atherosclerosis, nanoparticle–enabled in vivo RNAi mitigated blood leukocytosis, limited inflammation in the ischemic heart, and reduced myeloid cell numbers in atherosclerotic plaques.
understood, especially compared with the well-studied bone marrow niche. Understanding HSC retention factors and their regulation in the spleen was the purpose of this study.

Because the spleen harbors very few HSCs in the steady state, we investigated the splenic hematopoietic niche after injecting the Toll-like receptor ligand LPS to activate extra-medullary hematopoiesis. In the bone marrow, macrophages are an integral part of the HSC niche (Winkler et al., 2010; Chow et al., 2011) and differentiation depends on the receptor for macrophage colony-stimulating factor (M-CSFR, CD115; Auffray et al., 2009). We thus hypothesized that splenic hematopoietic niche assembly also requires M-CSFR signaling. In line with knockout studies (Takahashi et al., 1994; Dai et al., 2002), in vivo knockdown of M-CSFR with nanoparticle-encapsulated siRNA reduced splenic macrophage numbers substantially. Interestingly, decreased macrophage numbers were associated with a reduction of splenic HSCs. Depleting macrophages with diphtheria toxin (DT) in CD169 iDTR mice reproduced the findings obtained with M-CSF–directed siRNA treatment, thereby indicating that macrophages have a key role in splenic HSC maintenance. To investigate how splenic macrophages retain HSCs, we measured changes in splenic expression of major bone marrow retention factors after M-CSFR silencing. Silencing M-CSFR selectively reduced splenic VCAM-1, and the adhesion molecule was primarily expressed by macrophages. Inhibiting macrophage expression of VCAM-1 with siRNA targeting this adhesion molecule reduced splenic HSC numbers. Finally, we found that M-CSFR and macrophage–directed VCAM-1 silencing in mice with atherosclerosis mitigated blood leukocytosis and dampened inflammation in atherosclerotic plaques and the infarcted myocardium. These data reveal the importance of VCAM-1 expression by splenic macrophages for extramedullary hematopoiesis and illustrate the therapeutic potential of RNAi as an anti-inflammatory that mutates emergency overproduction and provision of myeloid cells.

**RESULTS**

**M-CSFR (CD115) knockdown attenuates LPS-induced splenomegaly**

Several siRNA sequences targeting the M-CSF receptor were selected using in silico prediction methods (Dahlman et al., 2014) and tested in vitro at two concentrations in M-CSFR–expressing cells. The six best duplexes (Fig. 1 A) underwent further testing at seven dilutions (Fig. 1 B). The siRNA with the best in vitro silencing at low concentrations (siCD115; sense, cuAucuAACcuucucGGAGAdTsdT; anti-sense, UUCCGGAGAAAGUUGAGuADTsdT) was incorporated into lipidoid nanoparticles (Love et al., 2010) for in vivo studies. These nanoparticles exhibit particular efficiency for in vivo silencing in monocytes and macrophages after systemic delivery (Leuschner et al., 2011; Novobrantseva et al., 2012; Majnudar et al., 2013; Courties et al., 2014) and resemble siRNA delivery vehicles currently used in clinical studies (Coelho et al., 2013). siCD115 was tested for in vivo silencing in the spleen, which has the highest siRNA concentration after systemic injection of nanoparticle carriers (Leuschner et al., 2011). Because the adult steady-state spleen harbors only a few HSCs, we induced extramedullary hematopoiesis by treating mice with LPS. Mice then received injections of either siCD115 or control siRNA with an irrelevant sequence (siCON), both encapsulated in similar lipidoid delivery nanoparticles (Fig. 1 C). We found that two intravenous siCD115 injections reduced CD115 mRNA levels in splenic monocytes and macrophages by eightfold (Fig. 1 D) and led to a substantial (>75%) in vivo protein knockdown in leukocytes (Fig. 1 E). Although the treatment did not significantly change body weight (Fig. 1 F), LPS-induced splenomegaly (Fig. 1 G) and splenic cell content (Fig. 1 H) fell significantly after siCD115 treatment. Moreover, siCD115 treatment significantly reduced serum IFN-γ levels (Fig. 1 I). Consistently, CD115 knockdown reduced numbers of total blood monocytes and Ly-6chigh monocytes in LPS-challenged mice (Fig. 1, J and K).

**M-CSFR knockdown decreases splenic HSPC numbers**

Next, we enumerated splenic Lin− Sca-1+ ckit+ CD48− CD150+ HSCs, LSKs (Lin− Sca-1+ ckit+ cells), and granulocyte-macrophage progenitors (GMPs) (Lin− Sca-1+ ckit+ CD16/32high CD34high) in splenocytes (Fig. 2, A and B). The naive mouse spleen contained few hematopoietic stem and progenitor cells (HSCs, 666 ± 166; LSKs, 3,400 ± 1,030; GMPs, 1,300 ± 337 per spleen; Fig. 2, A and B). LPS treatment increased these numbers 30-, 39-, and 33-fold, respectively. In parallel to the observation that RNAi curbed LPS-induced splenomegaly, we found that siCD115 treatment decreased splenic HSC, LSK, and GMP numbers significantly (Fig. 2, A and B). Similarly, M-CSFR knockdown reduced splenic HSC numbers (Fig. 2 C). Because CD115 signaling contributes to macrophage survival (Shaposhnik et al., 2010), we investigated whether the receptor has a similar effect on HSCs. CD115 knockdown did not affect HSC and LSK apoptosis in the bone marrow (Fig. 2 D) and spleen (Fig. 2 E).

**Splenic niche maintenance requires M-CSFR signaling**

To explore how M-CSFR knockdown reduced splenic HSPC numbers, we assessed HSPC proliferation using an in vivo BrdU incorporation assay, as M-CSF may facilitate cell proliferation. However, BrdU incorporation by HSPCs did not differ among the treatment groups, indicating that siCD115 does not change HSPC proliferation rates (Fig. 3 A). Because BrdU may act as a mitogen (Takizawa et al., 2011), we also used a CFSE dilution assay to measure LSK proliferation (Fig. 3 B). For this assay, we sorted LSKs from CD45.1+ mice, labeled them with CFSE, and transferred them into LPS-primed CD45.2+ mice treated with either siCD115 or siCON. Transferred CFSE-labeled LSKs diluted the dye at similar rates in both treatment cohorts, indicating similar rates of LSK proliferation, but the total number of splenic BrdU+ HSPCs declined after siCD115 treatment, reflecting reduced HSPC retention in the splenic niche (Fig. 3 A).
MacDonald, 2012), we next enumerated splenic macrophages after silencing this receptor. Treatment with siCD115 substantially reduced the numbers of splenic macrophages (Fig. 4 A). Because macrophages contribute to HSPC maintenance in the bone marrow (Winkler et al., 2010; Chow et al., 2011), we sought to clarify whether and how these cells might regulate the splenic niche. Splenic macrophages express CD169 (Fig. 4 B) and can be efficiently depleted in CD169-iDTR mice (Miyake et al., 2007) via DT treatment (Fig. 4 C). Similar to the effects observed after M-CSFR silencing, this macrophage depletion significantly reduced numbers of splenic HSCs, LSKs, and GMPs (Fig. 4 D). Bone marrow HSC and LSK numbers also fell significantly after macrophage depletion (Fig. 4 E). In contrast, HSCs, LSKs, and GMPs significantly increased in the blood (Fig. 4 F), indicating HSPC mobilization into circulation in the absence of macrophages. Collectively, these data indicate that M-CSFR signaling contributes to the hematopoietic niche by inducing macrophage differentiation.

The observed lower splenic HSPC retention suggested that M–CSFR knockdown might influence the expression of HSPC retention factors. Indeed, siCD115 treatment significantly reduced mRNA levels of the major retention factors in the bone marrow, including CXCL12, vascular cell adhesion molecule-1 (VCAM-1), angiopoietin, and stem cell factor (SCF; Fig. 3 C). Yet in the spleen, VCAM-1 was the only retention factor significantly reduced by siCD115 treatment (Fig. 3 D). Reducing these retention factors may favor progenitor release into the blood (Winkler et al., 2010). Accordingly, we found that M–CSFR knockdown increased the numbers of circulating LSKs by threefold (Fig. 3 E), indicating increased HSPC departure from their niches in the absence of M–CSFR signaling.

**Macrophages’ role in the splenic niche**

Because M–CSFR regulates macrophage differentiation, survival, and proliferation (Auffray et al., 2009; Hume and MacDonald, 2012), we next enumerated splenic macrophages after silencing this receptor. Treatment with siCD115 substantially reduced the numbers of splenic macrophages (Fig. 4 A). Because macrophages contribute to HSPC maintenance in the bone marrow (Winkler et al., 2010; Chow et al., 2011), we sought to clarify whether and how these cells might regulate the splenic niche. Splenic macrophages express CD169 (Fig. 4 B) and can be efficiently depleted in CD169-iDTR mice (Miyake et al., 2007) via DT treatment (Fig. 4 C). Similar to the effects observed after M-CSFR silencing, this macrophage depletion significantly reduced numbers of splenic HSCs, LSKs, and GMPs (Fig. 4 D). Bone marrow HSC and LSK numbers also fell significantly after macrophage depletion (Fig. 4 E). In contrast, HSCs, LSKs, and GMPs significantly increased in the blood (Fig. 4 F), indicating HSPC mobilization into circulation in the absence of macrophages. Collectively, these data indicate that M–CSFR signaling contributes to the hematopoietic niche by inducing macrophage differentiation.

![Figure 1. M-CSFR (CD115) knockdown reduces LPS-induced splenomegaly.](A) In vitro screening for siRNA inhibitors of CD115 mRNA (n = 3 per group). (B) Concentration dependence of the six best siRNAs against CD115 mRNA (n = 3). (C) Experimental design. LPS-treated mice were injected with siRNA against CD115 (siCD115) or control siRNA with an irrelevant sequence (siCON), or left uninjected. (D) CD115 mRNA levels in FACs-isolated monocytes/macrophages (n = 4–6). (E) CD115 expression on monocytes/macrophages by flow cytometry (n = 6–8 per group). (F) Body (n = 3–5) and (G) spleen weight of naive and treated mice (n = 3–10). (H) Total splenocyte count of naive and treated mice (n = 4–9 per group). (I) Serum IFN-γ levels measured by ELISA (n = 3–8). Flow cytometric plots (J) and quantification (K) of monocytes and Ly-6chigh monocytes in blood after siRNA treatment (n = 3–5). Data were pooled from 2–5 independent experiments and are mean ± SEM. Significance was determined by Mann-Whitney test or one-way ANOVA. **, P < 0.01; ***, P < 0.001.
To investigate if splenic VCAM-1+ macrophages are monocyte-derived or sourced from local progenitors, we induced parabiosis of GFP+ and GFP− mice (Fig. 5 H). After 6 wk of parabiosis, very few (<1%) VCAM-1+ macrophages were derived from the parabiont, i.e., from circulatory monocytes (Fig. 5, H and I). These data indicate that splenic VCAM-1+ macrophages arise from tissue progenitors rather than the bone marrow, in line with previously published data on splenic macrophages (Yona et al., 2013).

Our data suggest that splenic macrophages retain progenitors via expression of the adhesion molecule VCAM-1. To test this hypothesis directly, we formulated siRNA that targets VCAM-1 within macrophage-avid lipidoid nanoparticles (sense, AcuGGGuuGAcuuucAGGudTsdT; anti-sense, ACC-UGAAAGUcAACCcAGUdTsdT). This treatment, which limited VCAM-1 protein concentrations in macrophages but not in endothelial cells (Fig. 6 A), reduced splenic progenitor retention (Fig. 6 B). In contrast, numbers of HSCs, LSKs, and GMPs increased significantly in the circulation (Fig. 6 C).

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indicating mobilization of these cells into the blood after VCAM-1 knockdown in macrophages. In the bone marrow, HSPC numbers did not change significantly after VCAM-1 knockdown in macrophages (Fig. 6 D), pointing to retention mechanisms that compensate for reduced VCAM-1 levels in the bone marrow. Reduced splenic HSC retention may indicating mobilization of these cells into the blood after VCAM-1 knockdown in macrophages. In the bone marrow, HSPC numbers did not change significantly after VCAM-1 knockdown in macrophages (Fig. 6 D), pointing to retention mechanisms that compensate for reduced VCAM-1 levels in the bone marrow. Reduced splenic HSC retention may
attenuate LPS-induced myelopoiesis. To test this hypothesis, we enumerated myeloid cells and monocytes in the blood (Fig. 6 E) and spleen (Fig. 6 F) and found that VCAM-1 knockdown significantly reduced myeloid cell and monocyte numbers. Collectively, these data demonstrate a requirement for VCAM-1 expression by splenic macrophages for maintaining the organ’s hematopoietic niche. VLA-4 is an integrin expressed by leukocytes that binds with VCAM-1 on activated endothelial cells (Elies et al., 1990), resulting in recruitment of inflammatory cells to sites of inflammation. Similarly, VCAM-1 expressed by splenic macrophages may bind with VLA-4 expressed by HSCs (Williams et al., 1991). To investigate an interaction of VCAM-1 with VLA-4 in this setting, we transferred GFP+ HSPCs into LPS-treated mice after VLA-4 neutralization.
Figure 5. Splenic HSPCs reside near VCAM-1+ macrophages. (A) Gating for endothelial cells. CD45.2−CD31highICAM-2high cells were considered as endothelial cells. (B) VCAM-1 expression on splenic endothelial cell and leukocytes by FACS. (C) Cluster of GFP+ cells in the splenic red pulp 3 d after GFP+ HSPC transfer into wild-type mouse. Bar, 15 µm. (D) Cluster of GFP+ cells (green) near VCAM-1+ macrophages (red). Bar, 20 µm. (E) GFP+ cells (green) in close contact with VCAM-1+ (red) macrophages (F4/80+, blue) in the spleen. Bar, 5 µm. (F) 3D image of interaction between a GFP+ cell and a VCAM-1+ macrophage. (G) Fraction of GFP+ cells within indicated distance from VCAM-1+ macrophages (n = 5). (H) We induced parabiosis of GFP+ and GFP− mice. After 6 wk of parabiosis, the spleens of GFP− wild-type mice were analyzed for chimerism (GFP+ cells) using flow cytometry (H) and immunofluorescence microscopy (I; n = 3). Bar, 10 µm. Data were pooled from two independent experiments. Data are mean ± SEM. Significance was determined by one-way ANOVA and Mann-Whitney test. *, P < 0.05; ***, P < 0.01.

Figure 6. M-CSFR knockdown reduces inflammation in atherosclerotic plaque

with an antibody. We found that this blocking antibody significantly decreased splenic HSPC retention (Fig. 6 G).

M-CSFR knockdown reduces inflammation in atherosclerotic plaque

Previous studies revealed that splenic myelopoiesis provides inflammatory monocytes to atherosclerotic plaque (Swirski et al., 2007; Murphy et al., 2011; Robbins et al., 2012) and that, after experimental MI, splenic leukocyte overproduction accelerates progression of atherosclerosis (Dutta et al., 2012). A recent clinical trial indicated that similar mechanisms may exist in humans; 18F-FDG PET/CT showed increased metabolic activity in the spleens of patients with acute coronary syndrome (Kim et al., 2014). We thus investigated...
whether blocking splenic leukocyte overproduction with siCD115, which we found to disrupt the hematopoietic niche, would reduce systemic inflammatory activity in mice with atherosclerosis. ApoE^−/− mice fed a high fat diet received either siCD115 or siCON, as outlined in Fig. 7 A. In these mice, M-CSFR knockdown reduced splenic HSC and LSK
In accordance with reduced monocyte accumulation, plaque size, and necrotic core area decreased, and fibrous cap thickness increased in aortic root lesions of siCD115-treated mice (Fig. 7 E).

Figure 7. CD115 knockdown reduces inflammation in atherosclerosis. (A) Experimental design. (B) Enumeration of LSKs and HSCs in the spleens of ApoE−/− mice after CD115 knockdown. Representative FACS plots show percentage of monocytes in the blood (C), and macrophages and monocytes in the aorta (D) in ApoE−/− mice treated with either control or CD115 siRNA. The bar graphs show levels of monocytes and Ly-6Chigh monocytes in the blood (C) and macrophages and monocytes in the aorta (D) after CD115 knockdown (n = 4–5 per group). (E) Representative images showing Masson staining of the aortic root. The bar graphs depict quantification of total plaque size, fibrous cap thickness, and necrotic core area (n = 5 per group). Experiments were performed in duplicate. Bar, 30 µm. All data are shown as mean ± SEM. Statistical significance was determined by Mann-Whitney test. *, P < 0.05; **, P < 0.01.

Levels by 4- and 12-fold, respectively (Fig. 7 B), leading to much lower blood monocyte counts (Fig. 7 C). In the aorta, siCD115 treatment reduced the numbers of inflammatory myeloid cells, monocytes, and macrophages (Fig. 7 D).
M-CSFR and VCAM-1 knockdown reduces myocardial inflammation after ischemic injury

Coronary occlusion leads to inflammation in the infarcted myocardium (Swirski and Nahrendorf, 2013). To a large extent, infarct macrophages derive from the spleen (Swirski et al., 2009; Leuschner et al., 2012). Leukocyte oversupply or disturbed resolution of inflammation in the heart inhibits the infarct healing process, leading to aggravated left ventricular remodeling and heart failure (Nahrendorf et al., 2007; Panizzi et al., 2010). Although these preclinical observations likely apply to patients after MI (Tsujioka et al., 2009), no therapy targeting post-MI leukocyte oversupply to the heart has improved outcomes to date. We thus investigated whether silencing M-CSFR decreases inflammation in the ischemic heart. To this end, we treated mice with either siCD115 or siCON after coronary ligation (Fig. 8 A). Indeed, siCD115 treatment reduced numbers of myeloid cells, macrophages,

and monocytes in the infarcting myocardium 4 d after coronary ligation (Fig. 8 B). Likewise, histologically examined infarct tissue harvested on day 4 after coronary ligation showed significantly reduced Ly-6G and CD11b+ cells after siCD115 treatment (Fig. 8 C). Consistent with this, siCD115-treated mice had lower levels of myeloid cells and monocytes in the blood on day 4 after MI (Fig. 8 D).

Next, we investigated the effect of VCAM-1 knockdown in macrophages on day 4 after myocardial infarction. siVCAM-1 treatment significantly reduced myeloid cell, monocyte, and macrophage numbers in the infarct (Fig. 9, A and B). Of note, VCAM-1 knockdown did not alter proliferation and apoptosis of infarct macrophages (Fig. 9 C), indicating reduced supply of monocytes after the treatment. Consistent with this notion, we found significantly reduced myeloid cell and monocyte numbers in the blood (Fig. 9 D), bone marrow (Fig. 9 E), and spleen (Fig. 9 F).
levels of macrophage retention and egress cues netrin-1 (van Gils et al., 2012) and CCR7 (Trogan et al., 2006). VCAM-1 knockdown did not significantly change their expression (Fig. 10 F). This treatment reduced LSK and HSC numbers in the spleen (Fig. 10 G), thereby confirming that, in the setting of atherosclerosis, splenic macrophages retain HSCs via VCAM-1. Numbers of Ly-6c high monocytes in the spleen (Fig. 10 H) and blood (Fig. 10 I) fell. VCAM-1 silencing in macrophages significantly decreased plaque size and necrotic cores in the aortic root (Fig. 10 J). However, blood cholesterol levels were unchanged (Fig. 10 K).

**DISCUSSION**

Patients with myocardial infarction exhibit high leukocyte counts in the blood, inflamed arterial wall, and infarcted myocardium. Exaggerated inflammation in the heart, common in atherosclerotic individuals, promotes left ventricular dilation and heart failure. Extramedullary hematopoiesis, which contributes to leukocyte oversupply at sites of inflammation, occurs in systemic inflammatory conditions such as atherosclerosis (Murphy et al., 2011; Robbins et al., 2012) and after myocardial infarction (Leuschner et al., 2012). Hence, uncovering the pathways leading to extramedullary leukocyte production
as observed, for example, in the growing uterus of a pregnant mouse (Tagliani et al., 2011) and in peritoneal macrophages (Rosas et al., 2014). Moreover, M-CSF improves tissue macrophage survival by reducing apoptosis (Shaposhnik et al., 2010). During atherogenesis, vascular endothelial cells and smooth muscle cells can produce M-CSF (Clinton et al., 1992), and M-CSF–deficient op/op mice have dramatically reduced atherosclerotic plaques (Qiao et al., 1997; Rajavashisth et al., 1998). Our data imply that M-CSF signaling also plays a role in splenic leukocyte production. Because M-CSFR knockdown also affected the bone marrow, our data do not provide information on the relative contribution of the spleen versus bone marrow contribution to the inflammatory response after differentiation of monocytes to macrophages depends on M-CSFR signaling (Auffray et al., 2009). Local M-CSF signaling may also regulate tissue-resident macrophage proliferation, as observed, for example, in the growing uterus of a pregnant mouse (Tagliani et al., 2011) and in peritoneal macrophages (Rosas et al., 2014). Moreover, M-CSF improves tissue macrophage survival by reducing apoptosis (Shaposhnik et al., 2010). During atherogenesis, vascular endothelial cells and smooth muscle cells can produce M-CSF (Clinton et al., 1992), and M-CSF–deficient op/op mice have dramatically reduced atherosclerotic plaques (Qiao et al., 1997; Rajavashisth et al., 1998). Our data imply that M-CSF signaling also plays a role in splenic leukocyte production. Because M-CSFR knockdown also affected the bone marrow, our data do not provide information on the relative contribution of the spleen versus bone marrow contribution to the inflammatory response after...
MI or LPS challenge, but previous data suggest that the splenic contribution to the infarct myeloid population is substantial (Leuschner et al., 2012). Previous studies showed nanoparticle uptake was highest in macrophages but not strictly limited to these cells; therefore, the described mechanisms may not be exclusive. Because macrophage turnover in atherosclerotic plaque (Robbins et al., 2013) and in acute MI may be rapid (Leuschner et al., 2012), therapeutically targeting the cell’s supply by silencing M-CSFR may be a viable way to dampen inflammation in cardiovascular disease. Because human atheromata express M-CSFR, such an intervention might also directly attenuate inflammation at the plaque level (Salomon et al., 1992).

In accordance with mRNA array data on red pulp macrophages, we found these cells express high levels of VCAM-1 and are located in direct proximity to hematopoietic progenitor clusters. HSCs express the VCAM-1 ligand VLA-4 and rely on VCAM-1–VLA-4 interaction to anchor themselves in hematopoietic niches (Williams et al., 1991; Dutta et al., 2012). This type of cell–cell interaction is common in the spleen, which also retains marginal zone B cells (Lu and Cyster, 2002) and innate response activator B cells (Rauch et al., 2012). The latter protect against sepsis via GM-CSF production in the splenic red pulp. Interestingly, GM-CSF expands splenic monocyte production in the setting of atherosclerosis (Robbins et al., 2012; Wang et al., 2014). Our current study reveals that VCAM-1+ red pulp macrophages play a central role in maintaining splenic hematopoiesis.

Early after ischemic injury, increased angiotensin-II concentrations mobilize cells from the splenic monocyte reservoir, which contributes ~50% of leukocytes to the infarct. Later, the spleen continues to supply myeloid cells to the ischemic as well as the remote myocardium, now relying on IL-1β and SCF signaling to enhance splenic monocyte production (Dutta et al., 2012). Hematopoietic activity in the spleen follows the recruitment of bone marrow HSPCs, which circulate in higher numbers after MI in mice (Dutta et al., 2012) and in patients (Massa et al., 2005; Assmus et al., 2012). The relocation of HSPCs from the murine bone marrow to the spleen accelerates production of inflammatory myeloid cells, exacerbating preexisting atherosclerosis (Dutta et al., 2012). HSPC retention in hematopoietic tissue in the bone marrow and the spleen (Williams et al., 1991; Scott et al., 2003; Dutta et al., 2012) relies at least partially on the integrin VLA-4, a ligand for VCAM-1, which we report to be expressed by splenic macrophages. Even without an acute inflammatory trigger, such as MI, splenic myelopoiesis occurs in mice with chronic atherosclerosis (Robbins et al., 2012). In ApoE−/− mice on a high fat diet, HSPCs differentiate into monocytes in the spleen, a process that depends on GM-CSF and IL-3 (Wang et al., 2014). These monocytes then enter the circulation and migrate to atherosclerotic plaques. In the vessel wall, the cells and their progeny secrete inflammatory cytokines, give rise to reactive oxygen species, and produce proteases, all of which may render atherosclerotic plaques vulnerable to rupture (Moore and Tabas, 2011).

Despite growing evidence of the importance of inflammation in atherosclerosis and heart failure, specific antiinflammatory therapy has yet to be implemented in the clinic. Ongoing trials explore whether antiinflammatory therapeutics, such as anti–IL1β–neutralizing antibodies (CANTOS, NCT01327846) and methotrexate (CIRT, NCT01594333) reduce ischemic complications and death in patients with atherosclerosis. In addition to biologicals or small molecule drugs, RNAi may therapeutically interfere with the life cycles of inflammatory leukocytes that drive atherosclerosis progression and complication. Recent progress in RNAi delivery has led to clinical trials (Coelho et al., 2013) and enabled leukocyte silencing in rodents and primates (Novobrantseva et al., 2012). The 70-nm particles used in the current study of siRNA delivery to macrophages were identified in large-scale screens (Love et al., 2010), are rapidly synthesized from four components, and are similar to safe materials advancing through clinical trials. Our data indicate that RNAi targeting of hematopoiesis could reduce systemic inflammation by curtailing leukocyte overproduction. Silencing M-CSFR expression disrupted the splenic hematopoietic niche assembly, potentially allowing for decisive antiinflammatory intervention in individuals with vulnerable plaques or shortly after myocardial infarction. Currently, this approach would lend itself to short-term, systemic, parenteral delivery rather than sustained oral therapy. Longer therapeutic courses might compromise the salutary functions of M-CSF signaling and macrophages in host defenses. Because the siRNA delivery in our study was systemic and therefore not limited to splenic cells, we are unable to define to what precise degree the observed therapeutic effects derive from silencing in the spleen. Likely, monocyte differentiation in the bone marrow or at the sites of inflammation, i.e., the heart and the arterial wall, also contributed to lower macrophage numbers. Yet, the parallel and profound reduction of monocytes in the circulation and inflamed tissue indicates that lowering the systemic supply of inflammatory cells played a major role in reducing inflammation in the target tissue.

In conclusion, this study identifies a novel role for signaling through the M-CSFR in the genesis of splenic myelopoiesis. Interfering with this receptor inhibits macrophage functions in the splenic hematopoietic niche. These cells express VCAM-1 to retain HSPCs in the splenic red pulp. Interrupting this pathway with myeloid–directed in vivo RNAi, by silencing either M-CSFR or VCAM-1, enables HSPCs to escape from niches into blood and substantially reduces circulating monocytes. Ours is a first study on how RNAi affects hematopoiesis. Finally, this intervention decreases inflammatory cell numbers in atherosclerotic plaque and in the ischemic myocar-}

**MATERIALS AND METHODS**

**Animal strains and surgical procedures.** C57BL/6J, B6. SJL-Ptprc Pepcb/BoyJ (CD45.1), and B6.129P2-Apoetm1Unc/J (ApoE−/−) mice were purchased from The Jackson Laboratory. ApoE−/− mice consumed a...
high-cholesterol diet (Harlan Teklad; 0.2% total cholesterol) for 10 wk before the experiments. CD169−iDTR mice were gifts from M. Tanaka (Institute of Physical and Chemical Research Center for Allergy and Immunology, Yokohama, Japan). CD169−DTR litter mates were used as control. For other strains, appropriate control mice were purchased from Jackson Laboratory. All animal experiments were approved by Massachusetts General Hospital’s Institutional Animal Care and Use Committee.

To induce myocardial infarction, mice were intubated and ventilated with 2% isoflurane, and then underwent thoracotomy in the fourth left intercostal space. The left coronary artery was permanently ligated with a monofilament nylon 8-0 suture. The thorax was closed with a 5-0 suture.

**siRNA formulation and in-vitro screening.** siRNAs duplexes were generated to target the CSFRI (M-CSFR) transcript. Single-strand RNAs were designed and produced at Ahylam Pharmaceuticals as described previously (Frank-Kamenetsky et al., 2008). The NIH 3T3 cell line was transfected with siRNAs targeting M-CSFR or nontargeting control siRNA complexed with Lipofectamine RNAiMAX Transfection Reagent (Invitrogen) at 0.5 and 5 nM final concentrations. Cells were harvested 24 h after transfection, and expression levels of M-CSFR mRNA were quantified by the branched DNA QuantGene 2.0 assay (Affymetrix) using mouse M-CSFR–specific probes. Dose response was performed on the best siRNA duplexes with an siRNA, with the best siRNA having an EC50 of 10 pM. This M-CSFR–targeted siRNA (scD115) was scaled up for in vivo studies and has the following sequence: cuAcuAAcuucucucGAAAdTdT (sense), UUCGGAGAAAGUUGAGuAGdTsdT (antisense), wherein lower case letters identify 2’OMe modified nucleotides. VCAM-1–targeting siRNA was identified similarly, except C2C12 cells were used for siRNA screening and qRT-PCR using TaqMan primers, and probes were mRNA quantification. The VCAM-1–specific duplex selected for in vivo studies has the following sequence: AcuGGGcuAcuucucGAGuAdTdT (sense), ACCUGAAAGUUCAACcuAGuAdTdT (antisense). M-CSFR and VCAM1 targeting siRNAs were independently encapsulated into nanoparticle formulations as described previously (Akan et al., 2008; Leuschner et al., 2011).

**Treatment protocol.** C57BL/6 mice were treated with 10 µg of LPS for three consecutive days before tissue harvesting on the fourth day. siRNA against either M-CSFR or VCAM-1 was administered i.v. at 0.5 mg/kg body weight on the first and third day. To test efficacy of the siRNA after myocardial infarction, siRNA against either M-CSFR or VCAM-1 was injected i.v. at 0.5 mg/kg body weight on the day of MI and 2 d after MI. ApoE−/− mice were injected with the siRNA three times per week for 3 wk. CD169−DTR mice were injected with DT i.v. at 10 µg/kg body weight on the first and fourth days before analysis on the seventh day.

**Organ harvesting and tissue processing.** Cardiac puncture was performed to draw blood with an Eppendorf tube without anticoagulant. Blood was collected via cardiac puncture in Eppendorf tubes without anticoagulant. Blood was collected via cardiac puncture in Eppendorf tubes without anticoagulant. Blood was col-

**Flow cytometry.** Single cell suspensions (300 µl in FACS buffer) were stained with a cocktail of antibodies against lineage markers (CD90 [clone 53–2.1], B220 [clone RA3–6B2], CD49b [clone DX5], NK1.1 [clone PK156], Ly-6G [clone IAB8], and Ter-119 [clone TER-119]). After 30 min of staining, the cells were washed in FACS buffer and passed through 40-µm filters, followed by red blood cell lysis in the filtrate. Fenums were flushed with ice-cold FACS buffer.

For hematopoietic stem and progenitor cell staining, we used biotin-conjugated antibodies against lineage markers CD11b (clone M1/70), CD11c (clone N418), and IL7rα (clone A7R34) in addition to the lineage antibodies used for leukocyte staining. This was followed by a secondary staining with antibodies against c-KIT (clone 2B8), Sca-1 (clone D7), CD16/32 (clone 2.4G2), CD34 (clone RAM34), CD115 (clone AFS98), CD150 (9D1), and CD48 (HM48-1).

**HSC proliferation assays.** Mice were injected with BrdU 24 h before organ collection, and intracellular BrdU staining was performed using BrdU flow kits (BD). For CFSE dilution assay, sorted LSKs were stained with CFSE (Invitrogen) according to manufacturer’s protocol and injected into B6 mice treated with LPS and either siCON or siCD115. BrdU incorporation and CFSE dilution in hematopoietic stem and progenitor cells were measured by an LSRII.

**HSC apoptosis assay.** 5,000,000 bone marrow cells or splenocytes were cultured in 1 ml of DMEM and cultured in a 38 cm dish for 3 h to induce apoptosis. Apoptotic HSCs were stained with Annexin V/dead cell apoptosis kit (Invitrogen) and quantified by flow cytometry.

**VLA-4 neutralization.** Mice were injected i.v. with 200 µg of either anti-mouse VLA-4 (BioXCell) or control rat IgG2b (BioXCell) 2 d after LPS injection. About 300,000 lineage c−c− cells sorted from GFP+ mice were adoptively transferred i.v. on the day of VLA-4 injection.

**Total blood cholesterol measurement and ELISA.** Blood was collected via cardiac puncture in Eppendorf tubes without anticoagulant. Blood was kept at room temperature for ~2 h and spun down at 4°C for 10 min to separate serum. Serum was stored at −80°C, and total cholesterol was measured using an enzymatic colorimetric assay (Cholesterol E; Wako). We used Quantikine Colorimetric Sandwich ELISA kit (R&D Systems) to measure serum IFN-γ levels.

**FACS.** Bones were harvested, including all long bones and the spine. Single-cell suspensions were prepared by grinding bones with a pestle and mortar and passing the cells through 40-µm filters. The cells were stained with biotinylated antibodies against lineage markers for hematopoietic stem and progenitor cells, as described above, and then incubated with streptavidin MicroBeads (Miltenyi Biotec). Lineage− cells were depleted using MACS columns (Miltenyi Biotec). Lin+ cells were stained with antibodies for hematopoietic stem and progenitor cells as described above. LSKs were sorted using a FACSAria Ilu cell sorter (BD).

**RNA extraction and quantification.** RNA was extracted using PicoPure RNA isolation kit (Applied Biosystems) and eluted in a total volume of 10 µl. RNA was quantified using Nanodrop, and one microgram of mRNA was used to generate complimentary DNA (cDNA) using a high capacity RNA to cDNA kit (Applied Biosystems). Target genes were quantified using TaqMan gene expression assays (Applied Biosystems).

**Histology.** For histological analysis, aortas and hearts were embedded in OCT compound (Sakura Fineteck), and fresh-frozen serial 6 µm thick sections were prepared. Mason Trichrome staining was performed according to the manufacturer’s instructions (Sigma-Aldrich) to assess plaque size, fibrous cap thickness and necrotic core. Immunohistochemistry was performed using Ly-6G (IA8, BioLegend) and CD11b (M1/70, BD) antibodies, biotinylated anti-rat IgG antibody (Vector Laboratories Inc.) and VECAStain ABC kit (Vector Laboratories). The reaction was visualized using a 3-amino-9-ethylcarbazole (AEC) substrate (Dako), and all sections were counterstained with Harris hematoxylin solution (Sigma-Aldrich). The slides were scanned using Nanozoomer 2.0 RS (Hamamatsu), and staining was quantified in 5 high power fields per mouse, using semiautomated thresholding in IPlab, and expressed as percentage of positive area.
Immunofluorescence microscopy. 3 d after priming with LPS, the spleen was harvested and fixed by immersion in a 4% PFA PBS solution. Fixed tissues were cryoprotected in a 30% sucrose, 0.02% sodium azide, PBS solution at 4°C. Tissues were embedded and frozen in tissue freezing media (Leica) and then sectioned in a −20°C cryostat (Leica) at a thickness of 10 and 25 µm.

The tissue sections were stained with an anti–VCAM-1 antibody (sc-1504; Santa Cruz Biotechnology) and an anti-F4/80 (14–4801; eBioscience) antibody followed by a Cy3 donkey anti–rabbit and Cy5 donkey anti–rat IgG (Jackson Immunoresearch Laboratories). Sections were mounted in Vectashield + DAPI mounting media.

For Ki-67 immunofluorescence imaging, heart tissue sections were stained using an anti–Ki67 antibody followed by a Cy3-conjugated donkey anti–rabbit antibody. Sections were mounted in Vectorshield with DAPI mounting media. Digital image files were post-processed with Nikon Imaging Software Elements, Volocity 6 (Perkin Elmer), ImageJ, and IP laboratory.

We thank Dr. Masato Tanaka (Institute of Physical and Chemical Research Center for Allergy and Immunology, Yokohama, Kanagawa, Japan) for providing CD169+ DTR mice and the Alnylam siRNA synthesis and formulations groups for siRNA efficacy of RNAi therapy for transthyretin amyloidosis. We thank Dr. Masato Tanaka (Institute of Physical and Chemical Research Center for Allergy and Immunology, Yokohama, Kanagawa, Japan) for providing CD169+ DTR mice and the Alnylam siRNA synthesis and formulations groups for siRNA efficacy of RNAi therapy for transthyretin amyloidosis.

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