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Claire E. Olingy, Georgia Institute of Technology
Cheryl L. San Emeterio, Georgia Institute of Technology
Molly E. Ogle, Georgia Institute of Technology
Jack R. Krieger, Georgia Institute of Technology
Anthony C. Bruce, University of Virginia
David D. Pfau, Georgia Institute of Technology
Brett T. Jordan, Georgia Institute of Technology
Shayne M. Peirce, Georgia Institute of Technology
Edward Botchwey, Emory University

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Non-classical monocytes are biased progenitors of wound healing macrophages during soft tissue injury

Claire E. Olingy1, Cheryl L. San Emeterio1, Molly E. Ogle1, Jack R. Krieger1, Anthony C. Bruce2, David D. Pfau1, Brett T. Jordan1, Shayn M. Peirce1 & Edward A. Botchwey1

Successful tissue repair requires the activities of myeloid cells such as monocytes and macrophages that guide the progression of inflammation and healing outcome. Immunoregenerative materials leverage the function of endogenous immune cells to orchestrate complex mechanisms of repair; however, a deeper understanding of innate immune cell function in inflamed tissues and their subsequent interactions with implanted materials is necessary to guide the design of these materials. Blood monocytes exist in two primary subpopulations, characterized as classical inflammatory or non-classical. While classical monocytes extravasate into inflamed tissue and give rise to macrophages or dendritic cells, the recruitment kinetics and functional role of non-classical monocytes remains unclear. Here, we demonstrate that circulating non-classical monocytes are directly recruited to polymer films within skin injuries, where they home to a perivascular niche and generate alternatively activated, wound healing macrophages. Selective labeling of blood monocyte subsets indicates that non-classical monocytes are biased progenitors of alternatively activated macrophages. On-site delivery of the immunomodulatory small molecule FTY720 recruits S1PR3-expressing non-classical monocytes that support vascular remodeling after injury. These results elucidate a previously unknown role for blood-derived non-classical monocytes as contributors to alternatively activated macrophages, highlighting them as key regulators of inflammatory response and regenerative outcome.

The mononuclear phagocyte system plays a multi-faceted role in maintaining tissue homeostasis and responding to pathological processes such as autoimmune diseases, cancer, and aberrant wound healing. Monocytes circulate in the bloodstream during steady state and are robustly recruited to sites of inflammation, where they exert functions that include clearance of cellular debris, promotion of angiogenesis, and restoration of tissue integrity. The ontogeny of macrophages varies in different tissues, such that some tissue resident macrophages are seeded embryonically and self-renew in a similar manner to stem cells, whereas other macrophages (such as in the dermis or gut) are continually replenished by blood-derived monocytes. Consequently, circulating blood monocytes are considered a highly plastic and dynamic system of innate immune cells that initiate processes of organ and tissue remodeling. Immunologically smart interventions that exploit the division of labor between different monocyte and macrophage populations require an understanding of the roles that these cells play in promoting repair, as unchecked activity of innate immune cells can perpetuate tissue damage through chronic inflammation and fibrosis.

Two distinct subpopulations of monocytes have been identified in mouse and human blood, which can be distinguished by well-characterized surface protein expression profiles. Classical inflammatory monocytes are identified by Ly6C<sup>hi</sup>CX3CR1<sup>lo</sup>CD43<sup>lo</sup> expression in mice (CD14<sup>hi</sup>CD16<sup>−</sup> in human), whereas non-classical alternative monocytes are Ly6C<sup>lo</sup>CX3CR1<sup>hi</sup>CD43<sup>hi</sup> in mice (CD14<sup>+</sup>CD16<sup>+</sup> in human). A third population of intermediate monocytes characterized by intermediate expression of Ly6C in mice (CD14<sup>hi</sup>CD16<sup>+</sup> in humans) are...
thought to complement the functions of non-classical monocytes and may preferentially differentiate into dendritic cells within inflamed tissues\textsuperscript{9,10}. Under homeostasis, classical monocytes in blood decrease Ly6C expression and become non-classical Ly6C\textsuperscript{lo} monocytes\textsuperscript{11}, which patrol the luminal side of resting endothelium\textsuperscript{12}. Classical monocytes also survey steady-state tissues and can traffic to lymph nodes without differentiating into macrophages\textsuperscript{13}. During inflammation, monocytes exit peripheral blood and extravasate into tissue, where they may transiently persist as monocytes without differentiation and exert a host of functions within the damaged tissue\textsuperscript{13-17}. Classical inflammatory monocytes present in the acute phases of injury secrete pro-inflammatory cytokines such as IL-6, iNOS, and TNFα\textsuperscript{18} and exhibit high levels of matrix metalloproteinase and cathepsin production\textsuperscript{18}. Conversely, Ly6C\textsuperscript{hi} monocytes present later during inflammation secrete high levels of vascular endothelial growth factor (VEGF) and IL-10 and can induce endothelial cell proliferation to promote arteriogenesis\textsuperscript{14,18,19}. We have previously shown that strategies that enhance the early recruitment of Ly6C\textsuperscript{lo} monocytes couple with later increases in arteriolar expansion and angiogenic activity\textsuperscript{20,21}. Recruited monocytes can differentiate into macrophages, serving as an alternative source of wound macrophages to those derived from in situ proliferation of tissue resident populations\textsuperscript{22}.

Macrophages are highly responsive to cues within the injury niche, enabling them to dynamically modify their behavior in response to changes in the microenvironment and display extremely varied phenotypes. Classically activated (“M1”) macrophages are primary players in pathogen destruction, secretion of inflammatory cytokines, and driving Th1-type responses\textsuperscript{23}. Conversely, alternatively activated wound healing (“M2”) macrophages (of which a number of subtypes have been described\textsuperscript{23}) are associated with pro-regenerative activities such as angiogenesis\textsuperscript{24,25}, extracellular matrix remodeling\textsuperscript{26}, secretion of anti-inflammatory cytokines\textsuperscript{27}, and resolution of inflammation\textsuperscript{28}. The highly complex and heterogeneous nature of inflamed tissue microenvironments has rendered a general description of macrophage origin and function challenging. Within toxin-induced muscle injury\textsuperscript{29}, liver fibrosis\textsuperscript{30}, infection\textsuperscript{31}, and autoimmune disease\textsuperscript{32}, classical Ly6C\textsuperscript{lo} monocytes are recruited from circulation and undergo in situ differentiation to be the primary contributors of injury Ly6C\textsuperscript{lo} macrophages. In contrast, sequential recruitment of classical Ly6C\textsuperscript{hi} followed by non-classical Ly6C\textsuperscript{lo} monocyte subsets after myocardial infarction\textsuperscript{18}, and direct recruitment of adoptively transferred Ly6C\textsuperscript{lo} monocytes within excisional skin injury\textsuperscript{20} and during the development of inflammatory arthritis\textsuperscript{31} have been reported. However, whether specific populations of blood monocytes give rise to defined macrophage phenotypes surrounding implanted materials remains unknown.

Harnessing myeloid cell functions for regenerative medicine applications requires an understanding of the cues that direct the localization and fate of these cells. Immunoregenerative materials seek to leverage the function of endogenous immune cells to guide the progression of inflammation and repair damaged tissue\textsuperscript{32}. For example, local delivery of stromal-derived factor-1 (SDF-1) from desulfated heparin-containing poly(ethylene glycol) (PEG) hydrogels increases the frequency of CXCR4\textsuperscript{hi}Ly6C\textsuperscript{lo} monocytes, which promotes capillary network expansion\textsuperscript{31}. Moreover, co-delivery of macrophage colony-stimulating factor (M-CSF) with VEGF from PEG hydrogels increases the density and maturity of corneal blood vessels compared to VEGF alone\textsuperscript{33}. Recently, our lab demonstrated that poly(lactic-co-glycolic acid) (PLGA)-based delivery of the small molecule FTY720, an agonist of sphingosine-1-phosphate receptor (S1PR) 3, recruits non-classical monocytes to inflamed tissues and promotes arteriogenesis\textsuperscript{25}. In the present study, we utilize cell labeling strategies to selectively track the fate of either classical Ly6C\textsuperscript{hi} or non-classical Ly6C\textsuperscript{lo} monocytes in response to biomaterial implantation within cutaneous wounds. We demonstrate that classical monocytes are able to give rise to both CD206\textsuperscript{+} and CD206\textsuperscript{-} macrophages following monocyte depletion with clodronate liposomes, but labeled non-classical monocytes preferentially give rise to CD206\textsuperscript{-} M2-like macrophages. On-site delivery of the immunomodulatory small molecule FTY720 induces homing of extravasated non-classical monocytes to peri-implant vasculature. Subsequently, FTY720 promotes in situ generation of wound healing macrophages and vascular remodeling within ischemic skeletal muscle. These results shed light on the fate of specific monocyte populations following biomaterial implantation after injury and indicate that non-classical monocytes are a promising therapeutic target for harnessing pro-regenerative inflammation to promote repair.

Results

Skin wounding and biomaterial implantation induces monocyte trafficking. The dorsal skinfold window chamber (DWC) model is a partial thickness excisional skin injury model that involves removing the epidermis and dermis to reveal the underlying sub-reticular vasculature. We have previously used this model to investigate the recruitment of distinct monocyte subsets to inflamed tissue surrounding biomaterial implants\textsuperscript{20,21,34}. In this study, we tracked the fate of monocytes that are recruited in response to cutaneous wounding and material implantation. DWC surgery and implantation of a polymeric poly(lactic-co-glycolic acid) (PLGA) thin film 1mm in diameter decreased the frequency of blood monocytes by nearly four-fold 1 day post-injury, followed by a three-fold elevation by 3 days post-injury compared to blood taken at day 0 prior to surgery (Fig. 1a,b). Monocytes were identified based on CD11b\textsuperscript{+} and SSC\textsuperscript{lo} expression, as Gr\textsuperscript{1hi}Ly6C\textsuperscript{lo} cells are SSC\textsuperscript{lo} granulocytes (Supplemental Figure S1a). A three-fold decrease in the frequency of circulating classical Ly6C\textsuperscript{hi} monocytes was observed in the first day post-injury, followed by a five-fold increase by day 3 (Fig. 1b). Non-classical Ly6C\textsuperscript{lo} monocytes also initially decreased (by five-fold), but then increased by five-fold relative to day 0 (Fig. 1b) and were the most abundant type of monocyte in blood for the duration of the study (Supplemental Figure S1b). Ly6C\textsuperscript{lo} monocytes were similarly decreased at day 1 by three-fold fold following injury but returned to baseline levels by day 3. These changes in circulating myeloid cell populations were accompanied by corresponding relative changes in bone marrow cell populations. At 3 days post-injury, higher frequencies of total, Ly6C\textsuperscript{hi} and Ly6C\textsuperscript{lo} monocytes were detected in bone marrow, while the frequency of Ly6C\textsuperscript{lo} monocytes decreased (Fig. 1c). To further probe the fate of blood myeloid populations that were altered after injury, we utilized in situ labeling techniques to track blood monocytes.
Injured skin recruits Ly6C<sup>lo</sup> monocytes that give rise to CD206<sup>+</sup> wound macrophages. Circulating monocyte subsets were labeled in vivo prior to DWC surgery to facilitate cell tracking of each monocyte population as they entered inflamed dorsal tissue surrounding polymer implants. Non-classical Ly6C<sup>lo</sup> monocyte labeling was performed by intravenous administration of fluorescent latex beads (Fig. 2a), as previously described<sup>35, 36</sup>. Within two hours of intravascular administration, latex beads are phagocytosed and equally distribute within Ly6C<sup>lo</sup> and Ly6C<sup>hi</sup> blood monocyte populations. However, by 24 hours after injection, latex beads primarily label Ly6C<sup>lo</sup> monocytes due to physiological conversion of labeled Ly6C<sup>hi</sup> monocytes and this labeling is sustained for up to 1 week<sup>35</sup>. Utilizing this strategy, blood Ly6C<sup>lo</sup> monocytes were selectively labeled compared to classical Ly6C<sup>hi</sup> monocytes after DWC (83.4 ± 9.7% Ly6C<sup>lo</sup> monocytes vs. 14.5 ± 9.3% Ly6C<sup>int</sup> monocytes and 0.0 ± 0.0% Ly6C<sup>hi</sup> monocytes, day 1 post-injury) and the label was retained at similar proportions for the duration of the study (Fig. 2a, Supplemental Figure S1c,d). Analysis of digested explanted dorsal skin tissue (Supplemental Figure S2a,b) showed labeled cells originating from Ly6C<sup>lo</sup> monocytes primarily remained Ly6C<sup>lo</sup> within tissue. The lower frequency of Ly6C<sup>hi</sup> monocytes carrying the label in the tissue (9.6 ± 2.6% of LX + CD11b<sup>+</sup>SSC<sup>lo</sup> cells) compared to the frequency of labeled Ly6C<sup>lo</sup> cells (44.3 ± 8.2% of LX + CD11b<sup>+</sup>SSC<sup>lo</sup> cells) indicates that circulating Ly6C<sup>lo</sup> monocytes do not become Ly6C<sup>hi</sup> post-extravasation (Fig. 2b,c). Approximately half of labeled cells expressed F4/80, indicating that around half of recruited Ly6C<sup>lo</sup> monocytes convert into macrophages or are phagocytosed by macrophages by 3 days post-injury (Fig. 2d). About half of total F4/80<sup>+</sup>CD11b<sup>+</sup> cells expressed...
CD206 (Supplemental Figure S2c), whereas labeled cells were more likely to be immunophenotyped as CD206+ macrophages (91.0 ± 3.4% of LX+F4/80+CD11b+ cells) than CD206− macrophages (Fig. 2e). These data suggest that blood-derived Ly6Chi monocytes preferentially give rise to CD206+ wound repair macrophages in inflamed tissue surrounding material implants.

We then tracked circulating Ly6Clo blood monocytes to explore whether these cells adopted similar fates during inflammation. Ly6Clo monocytes were labeled by sequential administration of clodronate-loaded liposomes 2 days prior to injury followed by latex bead injection 16 h later (Fig. 2f). Clodronate liposomes administered intravascularly transiently deplete all blood monocytes, resulting in accumulation of latex beads in bone marrow cells that reappear in circulating Ly6Chi monocytes 2 days later35. Pre-administration of clodronate liposomes before latex bead injection preferentially labeled Ly6Chi monocytes (70.8 ± 13.3% Ly6Chi monocytes vs. 26.9 ± 12.9% Ly6Clo monocytes, day 1 post-injury; Fig. 2f). Labeled monocytes collected from digested dorsal tissue 3 days post-injury. (i) Frequency of bead-labeled F4/80+ cells in animals with labeled blood Ly6Chi monocytes. (j) Frequency of bead-labeled CD206+ cells out of total F4/80+ cells. Data presented as mean ± S.E.M. Statistical analyses performed using two-tailed t-tests. *p < 0.05, n = 4–11 animals per group.

**Figure 2.** Circulating non-classical Ly6Clo monocytes selectively give rise to CD206+ wound repair macrophages. (a) Latex (LX) beads were injected 1 day prior to DWC surgery to selectively label blood Ly6Clo monocytes. Right, flow cytometry dot plot of labeled blood LX+CD11b+SSClo cells shows selective Ly6Clo monocyte labeling 1 day post-injury. (b,c) Labeled monocytes collected from digested dorsal tissue 3 days post-injury. (d) Frequency of bead-labeled F4/80+ cells in animals with labeled blood Ly6Clo monocytes. (e) Frequency of bead-labeled CD206+ cells out of total F4/80+CD11b+ cells. (f) Blood Ly6Chi monocytes were selectively labeled by first depleting blood monocytes with i.v. clodronate liposome (Clod Lip) administration 2 days prior to injury, followed by LX bead-based labeling 16 h later. Right, flow cytometry dot plot of labeled blood LX+CD11b+SSClo cells shows selective Ly6Chi monocyte labeling 1 day post-injury. (g,h) Labeled monocytes collected from digested dorsal tissue 3 days post-injury. (i) Frequency of bead-labeled F4/80+ cells in animals with labeled blood Ly6Chi monocytes. (j) Frequency of bead-labeled CD206+ cells out of total F4/80+ cells. Data presented as mean ± S.E.M. Statistical analyses performed using two-tailed t-tests. *p < 0.05, n = 4–11 animals per group.
situ into Ly6C<sup>lo</sup> monocytes, as a significantly greater frequency of labeled Ly6C<sup>hi</sup> monocytes than Ly6C<sup>lo</sup> monocytes were detected (Fig. 2h). The frequency of F4/80<sup>+</sup> cells within the latex bead-positive (LX<sup>+</sup>) population was not different between Ly6C<sup>hi</sup> monocyte and Ly6C<sup>lo</sup> monocyte labeling (47.5 ± 17.3% vs. 48.0 ± 6.2% of LX<sup>+</sup> cells), suggesting that both monocyte populations are equally capable of acquiring a macrophage phenotype after extravasation (Fig. 2d,i). Labeled Ly6C<sup>hi</sup> monocytes showed no preference for acquiring CD206 expression within 3 days of injury (Fig. 2j), indicating that although blood-derived Ly6C<sup>hi</sup> monocytes can contribute to CD206<sup>+</sup> macrophages after clodronate liposome administration, they do so at a lower frequency than Ly6C<sup>lo</sup> monocytes.

Reduction of circulating Ly6C<sup>lo</sup> monocytes impairs CD206<sup>+</sup> macrophage generation. Intravascular administration of clodronate liposomes transiently depletes all circulating monocytes; however, because blood Ly6C<sup>lo</sup> monocytes are primarily derived from the conversion of Ly6C<sup>hi</sup> monocytes, there is a delay in the repopulation of circulating Ly6C<sup>lo</sup> monocytes<sup>5, 11</sup>. We employed this tool to examine how decreasing the quantity of circulating non-classical monocytes impacts the generation of CD206<sup>+</sup> macrophages during wound healing. As expected, a deficit in blood Ly6C<sup>lo</sup> monocytes, but not Ly6C<sup>hi</sup> monocytes, was observed 5 days after clodronate administration (Fig. 3a). No differences in either monocyte subtype (Fig. 3b), total F4/80<sup>+</sup> macrophages (Fig. 3c), or CD206<sup>+</sup> macrophages (Fig. 3d) were observed in digested tissue 3 days post-injury; however, a lower frequency of CD206<sup>+</sup> macrophages was observed (Fig. 3e). These findings suggest that circulating Ly6C<sup>lo</sup> monocytes are likely major contributors to the population of CD206<sup>+</sup> wound macrophages.

Adoptively transferred Ly6C<sup>lo</sup> monocytes preferentially differentiate into CD301b<sup>+</sup>CD206<sup>+</sup> macrophages. To complement in vivo bead-based labeling strategies and further investigate the role that blood monocytes play in macrophage generation during inflammation, we adoptively transferred sorted CD45.1<sup>+</sup>Ly6C<sup>lo</sup>CD43<sup>hi</sup> or Ly6C<sup>hi</sup>CD43<sup>lo</sup> monocytes into CD45.2 mice at the time of DWC surgery (Fig. 4a,b). While Ly6C<sup>lo</sup>CD43<sup>hi</sup> are a slightly more restricted population of Ly6C<sup>lo</sup> monocytes, these two populations primarily overlap and the same is true for Ly6C<sup>hi</sup>CD43<sup>lo</sup> monocytes (Supplemental Figure S2d). At 3 days post-injury, we assayed donor cells for expression of monocyte and macrophage markers (Fig. 4c, Supplemental Figure S3). A very low frequency of CD45.1<sup>+</sup>CD45.2<sup>+</sup> cells were detected in digested dorsal tissue (Fig. 4d, Supplemental Figure S4a,b), indicating that few donor cells are present 3 days post-injury. We detected a modest, but insignificant increase in the number of total donor cells (normalized to tissue mass) originating as Ly6C<sup>lo</sup>CD43<sup>hi</sup> relative to those adoptively transferred as Ly6C<sup>hi</sup>CD43<sup>lo</sup> (Fig. 4e). CD301b marks a population of macrophages that appear in the midphase of skin wound healing and are required for effective cutaneous repair<sup>37</sup>. Following adoptive transfer of Ly6C<sup>hi</sup> monocytes, we found that 80.7 ± 10.4% of donor-derived CD206<sup>+</sup>F4/80<sup>+</sup> macrophages were CD301b<sup>+</sup>. Similarly, following adoptive transfer of Ly6C<sup>lo</sup> monocytes, we found that 84.2 ± 6.4% of donor-derived...
CD206+F4/80+ macrophages were CD301b+. A greater frequency of donor-derived wound healing CD301b+CD206+F4/80+ macrophages was detected in animals receiving adoptively transferred Ly6CloCD43hi monocytes compared to those that received Ly6C hiCD43lo monocytes (Fig. 4f). While F4/80 is present on all macrophages, co-expression of CD64 and MerTK exclusively distinguishes macrophages from monocytes. These two populations significantly overlap, as 25.3±2.1% of total F4/80+CD11b+ cells are also CD64+MerTK+, but nearly all CD64+MerTK+ cells are F4/80+ (Fig. 4c, Supplemental Figure S4c) and 88.9±1.2% of CD206+F4/80+ cells are also CD64+MerTK+. Similarly, we detected a greater frequency of donor-derived CD301b+CD206+CD43 lo+MerTK+ macrophages in animals receiving adoptively transferred Ly6CloCD43 hi+ monocytes, supporting the conversion of these monocytes into pro-regenerative macrophages (Supplemental Figure S4f). No changes in the frequency of CD301b−CD206+ (F4/80+) (Fig. 4f), CD301b−CD206+CD64+MerTK−, or total number of macrophages (Supplemental Figure S4d,e,g) were detected between the two grafts. These results further support the hypothesis that circulating non-classical monocytes differentiate into wound repair macrophages.
On-site delivery of FTY720 promotes accumulation of alternatively activated macrophages.

We have previously demonstrated that localized delivery of the small molecule FTY720 from PLGA thin films enhances the recruitment of Ly6C<sup>lo</sup> monocytes to inflamed peri-implant tissue and supports arteriogenesis<sup>20,34</sup>. We explored the fate of specific blood monocyte populations in response to localized immune modulation by selectively labeling Ly6C<sup>lo</sup> and Ly6C<sup>hi</sup> monocytes followed by implantation of FTY720-loaded PLGA films in DWCs. Though statistically insignificant, a trend of increased frequency of blood-derived Ly6C<sup>lo</sup> monocytes was observed with on-site delivery of FTY720 compared to blank implant (Fig. 5a,c; Supplemental Figure S5a). FTY720 increased the frequency of CD206<sup>+</sup>F4/80<sup>+</sup>CD11b<sup>+</sup> macrophages within injured tissue 3 days post-surgery (Fig. 5b). The conversion efficiency of labeled blood-derived Ly6C<sup>lo</sup> monocytes into CD206<sup>+</sup> macrophages was similar between groups, suggesting that FTY720 does not enhance the rate at which monocytes convert to CD206<sup>+</sup> macrophages (Supplemental Figure S5b). Selective labeling of Ly6C<sup>lo</sup> monocytes demonstrates that FTY720 does not increase...
the frequency of circulation-derived Ly6C<sup>hi</sup> monocytes in tissue when Ly6C<sup>lo</sup> blood monocytes are reduced with clodronate liposomes (Fig. 5d,f). We observed no changes in the frequency of total CD206<sup>+</sup>F4/80<sup>+</sup>CD11b<sup>+</sup> cells (Fig. 5e), suggesting that FTY720 is unable to increase the number of alternatively activated macrophages after reduction of circulating Ly6C<sup>lo</sup> monocytes.

To investigate the molecular mechanisms of FTY720-mediated recruitment, we probed non-classical monocytes for expression of the sphingosine-1-phosphate receptor 3 (S1PR3), at which FTY720 exhibits agonist activity. We have previously demonstrated that FTY720 requires S1PR3 expression on hematopoietic cells in order to promote arteriogenic remodeling<sup>20</sup>. In addition to Ly6C, CX3CR1 (the fractalkine receptor) can be used to distinguish classical and non-classical blood monocytes<sup>39</sup>. While CX3CR1 is difficult to detect using antibody-based methods, CX3CR1<sup>GFP</sup>/+ transgenic mice enable real-time assessment of monocyte subset identity. CX3CR1<sup>hi</sup> monocytes primarily overlap with the Ly6C<sup>lo</sup> monocyte population, and conversely, CX3CR1<sup>lo</sup> monocytes are primarily Ly6C<sup>hi</sup> (Fig. 5g). S1PR3 surface expression is selectively higher on CX3CR1<sup>hi</sup> blood monocytes (Fig. 5h,i; Supplemental Figure S5c), which indicates that elevated S1PR3 expression is a signature of non-classical monocytes and is consistent with our previous studies that demonstrated higher S1PR3 mRNA and total protein in Ly6C<sup>lo</sup> monocytes<sup>20</sup>. Taken together, FTY720 likely increases the tissue content of CD206<sup>+</sup>macrophages by recruiting circulating S1PR3<sup>hi</sup> non-classical monocytes from circulation.

**CX3CR1<sup>hi</sup> monocytes localize to a perivascular niche in wounded skin.** Non-classical monocytes are arteriogenic and can promote re-vascularization of damaged tissue<sup>14, 20, 40</sup>. To investigate the spatial distribution and function of monocytes in inflamed tissue, we performed intravital confocal microscopy of tissue surrounding FTY720-loaded implants 1 day post-surgery in CX3CR1<sup>GFP</sup>/+ transgenic mice. CX3CR1<sup>hi</sup> and CX3CR1<sup>lo</sup> cells were identified based on fluorescent intensity (Supplemental Figure S5d). Delivery of FTY720 from implanted films increased the frequency of non-classical CX3CR1<sup>hi</sup> cells 1 day post-surgery (Fig. 6a) and significantly decreased the distance of CX3CR1<sup>hi</sup> cells, but not CX3CR1<sup>lo</sup> cells, to the nearest blood vessel (Fig. 6b–d). Additionally, CD68<sup>+</sup>CD206<sup>+</sup> macrophages visualized by immunofluorescence adopted an elongated morphology (as determined by possessing an aspect ratio larger than 2) along the vasculature surrounding FTY720 implants (Fig. 6e).

We have previously reported that FTY720 promotes expansion of peri-implant arterioles by 7 days post-injury<sup>20, 41</sup>. In the current studies, local delivery of FTY720 was able to induce modest expansion of arteriole microvessels (<50µm diameter) by 3 days post-injury (Supplemental Figure S6a–c). Reduction of circulating non-classical monocytes with clodronate liposomes impaired arteriogenic expansion of peri-implant vessels (Supplemental Figure S6d). Interestingly, FTY720 exhibited a negative impact on vessel expansion when Ly6C<sup>lo</sup> monocytes were...
reduced (Supplemental Figure S6d). Taken together, FTY720 recruits non-classical monocytes that convert into CD206+ macrophages that closely associate with peri-implant vasculature, where they support arteriogenesis.

To investigate whether perivascular localization is a signature of non-classical monocytes, we compared the spatial positioning of monocyte subsets in vitro and in vivo. CX3CR1hi and CX3CR1lo monocytes sorted from bone marrow of CX3CR1GFP/+ mice were co-cultured with murine endothelial cell networks on Matrigel. A significantly higher frequency of CX3CR1hi non-classical monocytes compared to CX3CR1lo classical monocytes localized to in vitro-forming vessel networks (Fig. 7a,b). To determine whether monocytes positioned themselves in proximity to the vessels, we compared the proportion of cells within 20 µm of the vessel to the positions of a computer-generated random distribution of cells. Random distributions were generated by stochastically positioning the same quantity of CX3CR1hi or CX3CR1lo monocytes on images of in vitro endothelial networks (Supplemental Figure S7). We were unable to distinguish randomly-generated distributions of cells from the experimental distribution of CX3CR1lo monocytes with respect to the proportion of cells in close proximity (less than 20 µm) to the endothelial network (Fig. 7b). Conversely, the frequency of CX3CR1hi monocytes in close proximity to the endothelial network was significantly higher than the random position distributions (Fig. 7b).

Intravital confocal imaging surrounding FTY720-loaded polymer implants indicated that CX3CR1hi cells were positioned closer to inflamed vasculature than CX3CR1lo cells (Fig. 7c–e). These results indicate that non-classical monocytes preferentially localize near endothelial cells, which likely enables them to exert their angiogenic and arteriogenic effects14, 25, 40.

FTY720 alters monocyte/macrophage accumulation and promotes vascular network expansion after arteriole ligation. The recruitment kinetics, fate, and function of myeloid cells during inflammation is heavily dependent on the specific type of tissue injury18, 28. Consequently, we sought to determine whether application of FTY720-loaded materials to ischemic muscle injury produces similar patterns of immunomodulation. Feeder arteriolar vessels within the murine spinotrapezius muscle were ligated in CX3CR1GFP/+ mice and unloaded or FTY720-loaded PLGA films were implanted over the muscle immediately after injury. We observed a decrease in the overall area of CX3CR1hi cells in FTY720-treated animals 3 days post-injury (Fig. 8a,b). Conversely, we observed more CD206+ cells (Fig. 8c) and the area ratio of CD206+ cells to CX3CR1hi cells was higher in FTY720-treated animals (Fig. 8d). Previous work has demonstrated that monocytes differentiate into alternatively activated macrophages in vascular niches42. We investigated the localization of CX3CR1hi and CD206+ cells with respect to lectin-perfused vasculature. Fewer CX3CR1hi cells (Fig. 8e), but more CD206+ cells
(Fig. 8f) were found within 50 μm of blood vessels in FTY720-treated animals, which is consistent with perivascular conversion of non-classical monocytes into alternatively activated macrophages. Additionally, FTY720-treated animals had a greater vessel density and total length of arterioles 3 days after ligation (Fig. 8g–i).

Discussion

Monocytes are bloodborne mononuclear phagocytes that support tissue homeostasis and exit the vasculature at increased rates to differentiate into macrophages and dendritic cells during inflammation. The precise relationship of circulating classical and non-classical monocyte subsets to defined macrophage populations remains unknown. We have shown that after skin wounding and biomaterial implantation, circulating non-classical S1PR3hi monocytes extravasate into inflamed tissue and serve as biased progenitors of CD206+CD301b+wound healing macrophages. Previous work has demonstrated that classical monocytes directly convert to non-classical monocytes and macrophages within inflamed tissue16, 19. Therefore, these studies elucidate a complementary role for non-classical monocytes and argue that these cells primarily differentiate into alternatively activated macrophages. Biomaterial-mediated strategies that increase recruitment of non-classical monocytes through S1PR signaling are a promising strategy to increase accumulation of alternatively activated, wound healing macrophages (Fig. 9).

Non-classical Ly6Clo monocytes patrol resting endothelium during homeostasis12, and are present in circulation at a slightly higher frequency than classical Ly6Chi monocytes (Fig. 1b, Supplemental Figure S1b; 54.8 ± 7.5% Ly6Clo monocytes vs. 30.4 ± 5.6% Ly6Chi monocytes out of CD11b+SSClo cells). Interestingly, we observed a transient reduction of circulating monocyte populations 1 day post-surgery, which could be due to acute cell recruitment from the blood to injured tissue, as monocytopoiesis occurs over several days43. As previously reported18, we observed elevation of circulating myeloid cells 3 days post-injury, including total CD11b+Ly6G−SSClo cells and Ly6Chi and Ly6Clo subpopulations (Fig. 1b). Though we detected a decrease in Ly6Cint monocytes 1 day post-injury, there was no elevation at 3 days post-injury, indicating that the size of this population may not be altered by inflammatory stimuli. The observed changes in circulating cell populations during inflammation are likely due in part to alterations in myeloid cell trafficking from the bone marrow, which displayed parallel changes in myeloid composition (Fig. 1c). We expect that these systemic changes in blood myeloid populations are a result of DWC surgery and not material implantation, as we have previously demonstrated that the number of rolling
and adherent CX3CR1+ cells within dorsal skin vasculature is not significantly different between animals undergoing only DWC surgery and those that also received a PLGA implant.

Intravascular non-classical Ly6Clo monocytes orchestrate the disposal of necrotic endothelial cells after activation with TLR7-targeted danger signals during inflammation, whereas extravascular Ly6Clo monocytes can promote angiogenesis and matrix remodeling via secretion of VEGF and matrix metalloproteinases. The unique protein signature of non-classical monocytes, characterized by higher VEGF, TGFβ, and IL-10, and lower TNFα and IL-1β expression compared to classical monocytes has resulted in the suggestion that these cells may constitute a class of “anti-inflammatory” monocytes. Previous studies have indicated that Ly6Clo monocytes in inflamed tissues such as skeletal muscle and focal hepatic injury are derived from cells recruited as Ly6CHi monocytes from the blood and are converted in situ to Ly6Clo monocytes to promote wound healing and tissue repair. Other reports suggest that Ly6Clo monocytes are directly and robustly recruited from the blood, leading to extravascular accumulation during inflammation. While we observed that blood-derived Ly6CHi monocytes enter inflamed tissue, reduce Ly6CHi expression (Fig. 2g), and differentiate into macrophages (Figs 2i and 4f), our studies indicate that Ly6Clo monocytes are also directly recruited from circulation to injured tissues and able to give rise to macrophages (Figs 2c and 4f). Previous studies have indicated that adoptively transferred Ly6CHiMHCII+ monocytes do not infiltrate the skin but this may be a result of differences in the grafted cell population (Ly6CHiMHCII+ monocytes vs. Ly6CloCD43hi monocytes in our studies) or inflammatory stimuli (LPS injection vs. excisional skin injury). Though we observed a decrease in Ly6Clo blood monocytes after clodronate liposome administration, we did not detect a difference in the frequency of Ly6Clo monocytes within injured skin. Many studies have shown that Ly6Clo monocytes can convert to Ly6Clo monocytes in the blood or tissue and these results may reflect that in the absence of Ly6Clo monocytes, circulation-derived Ly6Clo monocytes compensate and increase their rate of conversion into Ly6Clo monocytes in inflamed tissue.

An important finding of our work is that circulating Ly6Clo monocytes preferentially contribute to the CD206+ wound macrophage pool within skin injury compared to Ly6CHi monocytes. Analysis of peri-implant tissue for recruited latex bead-labeled Ly6Clo monocytes shows that this subset is predisposed to acquire a CD206+ alternatively activated M2-like macrophage phenotype (Figs 2e and 4f). Administration of clodronate liposomes enables selective labeling of Ly6CHi monocytes, while simultaneously reducing the frequency of Ly6Clo monocytes for 2–7 days after administration. Clodronate liposome-treated animals displayed no change in the frequency of overall macrophages within the skin injury after 3 days, but exhibited a decrease in the proportion of CD206+ macrophages (Fig. 3c,e). Because tissue monocyte composition was unaffected by clodronate liposomes, these results may reflect a delay in generation of CD206+ macrophages. We cannot exclude the possibility that transient monocyte depletion alters myeloid cell responses and recruitment during inflammation, as monocyte subsets are known to communicate with each other.

Latex bead-based labeling strategies are useful because they overcome many of the limitations of adoptive transfer, including ex vivo cell manipulation, pooling of donor cells, and low sensitivity due to poor cell recovery. Importantly, there is no evidence that these labeling techniques alter monocyte recruitment or systemic inflammation. However, our interpretation of these studies relies on retention of latex beads within the cell that was originally labeled and no transfer to other cells. Therefore, cell tracking is best performed by use of complimentary tracking methods, including adoptive transfer, selective labeling techniques, and transgenic or knockout mice. We adoptively transferred 5.55 × 10^6 CD45.1+ Ly6CHi or Ly6Clo monocytes by intravenous injection into CD45.2 mice at the time of DWC surgery. We did not detect differences in the total number of CD45.1+ donor-derived cells per milligram of dorsal tissue between the two monocytes grafts, and further probed for markers of macrophage differentiation and polarization within donor cells. CD301b+CD206+ macrophages drive midstage skin

Figure 9. Local FTY720 release recruits biased progenitors of wound healing macrophages to inflamed tissue. Circulating non-classical S1PR3+ monocytes are recruited by local delivery of FTY720 from a material implanted within an injury site. Upon entrance into inflamed tissue, non-classical monocytes give rise to alternatively activated, wound healing macrophages. Conversely, classical monocytes differentiate into both inflammatory and wound healing macrophages.
regeneration by promoting fibroblast repopulation, cellular proliferation, and re-epithelialization. In our adoptive transfer studies, a higher frequency of donor Ly6C hi monocytes acquired a CD301b+ CD206+ macrophage phenotype compared to donor Ly6C lo monocytes (Fig. 4f), indicating that circulation-derived Ly6C hi monocytes may be intrinsically predisposed to convert to dermal wound healing macrophages. Conversely, there was no difference in the frequency of donor-derived CD301b− CD206+ or number of total macrophages between mice receiving Ly6C hi or Ly6C lo monocytes (Fig. 4g, Supplemental Figure S4). These data, along with the observation that FTY720 administration increases CD206+ macrophages within injured skin (Fig. 5b), further supports the hypothesis that non-classical monocytes undergo local conversion into alternatively activated macrophages.

While macrophages are key mediators of the host response to implanted materials that govern the integration of implants into host tissue, the role that their monocytic precursors play in regulating implant outcome has been largely unexplored. In our studies, around 45% of monocytes surrounding PLGA implants were Ly6Clow/int. While lymphocyte, granulocyte, and macrophage accumulation are impacted by the type of material implanted, future studies are needed to determine how different classes of materials impact infiltration of monocyte and macrophage subsets. Interestingly, we observed model-specific differences in the kinetics of monocyte recruitment during FTY720 delivery. While FTY720 increases the frequency of Ly6C hi monocytes 3 days after skin wounding26 (Fig. 5a), we observed a decrease in CX3CR1hi monocytes 3 days after arteriole ligation in the spinotrapezius muscle (Fig. 5b). The progression of inflammation may be expedited during ischemia because of the need to rapidly restore blood supply and oxygen transport. FTY720 increases the frequency of F4/80+/ CD206+ macrophages both in wound skin (Fig. 5b) and CD68+/ CD206+ macrophages in ischemic muscle (Fig. 8f). Previously, we have detected more M2-like macrophages within mandibular bone defects 3 weeks after implantation of FTY720-loaded polymer scaffolds28. Taken together, these findings suggest that FTY720 delivery from different biomaterials can enhance pro-regenerative myeloid cell recruitment, but the kinetics may vary depending on the type of injury.

Monocytes and macrophages have been shown to interact closely with remodeling vasculature22. In vitro, CX3CR1hi monocytes preferentially associate with endothelial cell networks (Fig. 7a,b), and in vivo, exhibit closer homing to inflamed vasculature (Fig. 7c–e). While further studies are needed, it is possible that the vasculature provides microenvironmental cues that educate non-classical monocytes to convert into alternatively activated macrophages via signaling with the endothelium. Conversely, monocytes and macrophages appear to promote both angiogenic and arteriogenic expansion of the vasculature14, 52, 54. Macrophages are the primary source of angiogenic growth factors such as VEGF15 and robust vascularization is associated with increased macrophage presence in and around biomaterial implants13, 50. M2-polarized macrophages (IL-4– or IL-10-stimulated) are considered pro-angiogenic both in vitro and in vivo16–18, though recent studies have demonstrated that M1-polarized macrophages may also play key roles in angiogenesis19, 60. Promotion of both angiogenic expansion of vascular length and arteriogenic diameter network expansion in dorsal skin is correlated with the presence of Ly6C hi monocytes in higher proportion to Ly6C lo monocytes21. Hydrogel vascularization in response to growth factor delivery (platelet-derived growth factor and fibroblast growth factor) in the cornea was accompanied by accumulation of macrophages that produce both M1- and M2-associated mRNA transcripts, such as Tnfa and Arg1, respectively33.

We found that ischemic muscles treated with FTY720 have increased numbers of perivascular CD206+ cells located (Fig. 8f), which coincided with increased arteriolar length and total vessel density (Fig. 8g–i). FTY720 likely acts on both immune cells and the endothelium, as loss of S1PR3 in hematopoietic or parenchymal cells impairs FTY720-induced arteriogenic remodeling20. Delivery of FTY720 similarly induced both increased accumulation and perivascular localization of alternatively activated macrophages after volumetric muscle loss, which was accompanied by enhanced re-vascularization and muscle healing20. Consequently, both the phenotype of monocytes and macrophages, as well as their spatial distribution is likely an important feature when assessing their function.

Taken together, our studies shed light on the functions of non-classical monocytes during inflammation and wound healing. Monocytes and macrophages are increasingly appreciated for their roles in regulating tissue homeostasis and coordinating repair after damage. Acute modulation of the inflammatory response has been shown to regulate repair of tissue at longer time scales, including bone50 and skeletal muscle19. An understanding of the origin of monocyte and macrophage populations during wound healing and the cues governing their in situ fate is critical to harnessing endogenous mechanisms of repair. This work provides new insights into the origin of alternatively activated wound healing macrophages that can be leveraged to develop next-generation immunoregenerative biomaterials capable of finely tuning the inflammatory response.

Materials and Methods

Material fabrication. Films were fabricated as previously described20. Briefly, 350 mg PLGA (50:50 DLG 5E – Evonik Industries) was dissolved in 2ml dichloromethane in a glass scintillation vial via high-speed vortexing. For drug-loaded films, 1.75 mg of FTY720 (Cayman Chemical) was added at a 1:200 drug-polymer weight ratio, and mixed until completely incorporated. Polymer solutions were poured into Teflon-coated petri dishes and allowed to dry at −20 °C for 7 days. Before use, films were lyophilized overnight to remove any traces of solvent.

Dorsal skinfold window chamber surgery. All animal procedures were conducted according to protocols approved by the Georgia Institute of Technology or University of Virginia Institutional Animal Care and Use Committee. Male C57BL/6J or B6.129P-Cx3cr1tm1Litt/J mice (CX3CR1GFP/+ or CX3CR1GFP–/–) mice (8–12 weeks) were fitted with sterile dorsal skinfold window chambers (AP Trading Co) as previously described20. Briefly, mice were anesthetized with an intraperitoneal (i.p.) injection of ketamine/xylazine (100/10 mg/kg) in sterile saline. Dorsal skin was shaved, depilated, and sterilized with three alternating washes of 70% ethanol and chlorhexidine. A double-layered skin fold was elevated off the back of the mouse and fitted with the titanium frame of the window chamber on the underside. The epidermis and dermis were removed from the top side of the skinfold in a ~12 mm diameter circular area via surgical microscissors to reveal underlying vasculature. Exposed tissue was superfused
with sterile saline to prevent desiccation. The titanium frame was then mounted on the top side of the skinfold, attached to the underlying frame counterpart, and sutured to the surrounding tissue. Two films were placed on top of the exposed subreticular dermis layer immediately after surgery (day 0) and exposed tissue was sealed with a sterile glass window. Mice were euthanized 72 hours after surgery via CO₂ asphyxiation. The vasculature was immediately flushed with intracardiac infusion of saline followed by an intracardiac infusion of 4% paraformaldehyde, prior to whole mounting of tissue and immunohistochemistry.

**Spinotrapezius ischemia model.** Mice were anesthetized with an i.p. injection of ketamine/xylazine/atropine (60/40/2 mg/kg). Ligation surgeries were performed as previously described. Briefly, a small incision was made on the dorsum above the lateral edge of the right spinoptrapezius at the edge of the fat pad. The fascia was separated from the top of the muscle and the fat pad moved before isolating an anatomically reproducible feeding arteriole entering the muscle from below. This feeding arteriole was ligated with 10-0 nonabsorbable suture in two places and cut. The fat pad and fascia were moved back into position and the skin was closed with 8-0 nonabsorbable suture. To allow visualization of vascular endothelium of arterioles in CX3CR1-GFP mice, anesthetized mice were administered an intra-jugular injection of labeled isoelectric (IB4-Alexa Fluor 568; Life Technologies), which was allowed to circulate for 10 minutes. Anesthetized mice were euthanized via CO₂ asphyxiation 72 hours post-surgery. The vasculature was immediately flushed with an intracardiac infusion of adenosine (70 mg/L) in Ringer's solution followed by an intracardiac infusion of 4% paraformaldehyde.

**Flow cytometry.** Peripheral blood was collected via cardiac puncture and bone marrow was collected via centrifugation (1000 g for 5 mins) of isolated tibiae. The dorsal tissue circumscribing films was punched out with a 6 mm biopsy punch and pooled from both films within one animal for most studies. For adoptive transfer studies, all inflamed dorsal tissue was collected for analysis. Tissue was digested with collagenase (1 mg/ml) at 37 °C for 30 minutes and further disaggregated with a cell strainer to create a single cell suspension. Single cell suspensions of tissues were stained for flow cytometry analysis according to standard procedures and analyzed on a FACS-AriaIIIu flow cytometer (BD Biosciences). The following antibodies were used for cell phenotyping: APC-Cy7- or BV421-conjugated anti-CD11b (BioLegend), APC- or BV510-conjugated anti-Ly6C (BioLegend), PerCP-Cy5.5-conjugated anti-CD45 (BioLegend), PE-Cy7-conjugated anti-Gr-1 (BioLegend), APC-Cy7-conjugated anti-Ly6G (BioLegend), APC-conjugated anti-F4/80 (BioLegend), PE-Cy7- or FITC-conjugated anti-CD206 (BioLegend), PE-Cy7-conjugated anti-CD301b, BV711-conjugated anti-CD64 (BioLegend), PE-conjugated anti-MerTK (BioLegend), BV605-conjugated anti-CD45.1 (BioLegend), BV785-conjugated anti-CD45.2 (BioLegend), or PerCP-eFluor710 conjugated anti-CD115 (eBioscience). Dead cells were excluded by staining with Zombie NIR™ (Biolegend) in protein-free buffer prior to antibody staining. Staining using BV dyes was performed in the presence of Brilliant Stain Buffer (BD Biosciences). Positivity was determined by gating on fluorescence minus one controls. Absolute quantification of cell numbers in blood and tissue was performed by adding 25 μL of AccuCheck counting beads to flow cytometry samples (Thermo Fisher Scientific). S1PR3 flow cytometry was performed by first performing Fc block (Biolegend), followed by staining cells with primary unconjugated anti-S1PR3 antibody (Alomone Labs) and secondary staining with DyLight 650 anti-rabbit IgG (Abcam). Positivity was determined by staining CX3CR1-GFP+ cells with the secondary body only (no primary S1PR3 antibody).

**Cell tracking of Ly6C<sup>+</sup> and Ly6C<sup>hi</sup> monocytes.** For selective labeling of Ly6C<sup>+</sup> monocytes, mice were administered 250 μL of Fluoresbrite® Polychromatic Red latex beads intravenously one day prior to surgery (0.5 μm, Polysciences - diluted 1:25 in sterile saline) via jugular vein injection. For selective labeling of Ly6C<sup>hi</sup> monocytes, mice were administered 100 μL of clodronate liposomes per 10 g of mouse body weight (Dr. Nico van Rooijen, clodronateliposomes.com) intravenously two days prior to surgery, followed by administration of latex beads 16 hours later (one day prior to surgery). Labeling was confirmed by retro-orbital blood draw days 1 and 3 post-surgery. For adoptive transfer studies, white blood cells from bone marrow, spleen, and blood were collected from mice expressing the CD45.1<sup>+</sup> allele and enriched for monocytes using an EasySep Mouse Monocyte Isolation Kit (Stem Cell Technologies). Cells were further purified by fluorescence activated cell sorting on a BD FACS AriaIIIu flow cytometer and suspended in a SD FACSAriaIIIu cell sorter using the following markers: Ly6C<sup>+</sup> monocytes (SSC<sup>-</sup>CD11b<sup>+</sup>Ly6C<sup>hi</sup>CD45<sup>+</sup>CD43<sup>-</sup>) or Ly6C<sup>+</sup> monocytes (SSC<sup>-</sup>CD11b<sup>+</sup>Ly6C<sup>hi</sup>CD45<sup>+</sup>CD43<sup>-</sup>). Mice received 555,000 Ly6C<sup>+</sup> or Ly6C<sup>hi</sup> monocytes via jugular vein injection on the day of surgery.

**Intravitreal image acquisition.** Mice were anesthetized with isoflurane, the glass window was removed, and dorsal tissue was superimposed on saline to prevent desiccation. Up to two films were implanted into the window chambers on the day of surgery (day 0). To label perfused vasculature, mice were anesthetized with isoflurane and given a retro-orbital injection of high molecular weight TRITC-conjugated dextran (2 MDa; Life Technologies). For imaging, the anesthetized mouse was secured to the microscope stage in a custom adapter, the glass window was removed, and dorsal tissue was superfused with sterile saline. Intravital confocal microscopy was conducted with a 20X water immersion objective (NA = 1.0) on a Zeiss LSM710 NLO microscope and z-stack images were acquired immediately proximal to the films. For 3D analysis in Imaris (Bitplane), images of 708 × 708 μm regions were acquired adjacent to the implant to visualize immune cell distribution in the close surrounding tissue. Cells expressing CX3CR1-GFP were identified in Imaris using the surface tool. CX3CR1<sup>+</sup> surfaces were identified by smoothing with a 2 μm grain size and an automatic threshold on absolute intensity. Touching objects were split using a seed point diameter of 10 μm. CX3CR1<sup>hi</sup> versus CX3CR1<sup>+</sup> cells were discriminated by applying a filter to select surfaces with a high fluorescence intensity in the CX3CR1-GFP channel (above 150 max intensity). Vessels were identified in Imaris by drawing a surface on the TRITC-dextran fluorescent channel with a 3 μm grain size, manually-selected threshold value (determined based on each image), and manually-selected volume filter to remove small debris. To calculate the distance between CX3CR1<sup>+</sup> cells and the nearest blood vessel, a distance
transformation was applied to TRITC-dextran vessel surfaces and the median position of each CX3CR1⁺ cell within this space was recorded.

Whole mount immunohistochemistry. Dorsal tissue and spinotrapezius muscles were explanted and permeabilized overnight with 0.1–0.2% saponin. The tissues were blocked overnight in 5–10% mouse serum. Tissues were incubated at 4 °C overnight in solution containing 0.1% saponin, 5% mouse serum, 0.5% bovine serum albumin, and the following conjugated fluorescent antibodies: Alexa Fluor 594 anti-CD31 antibody (BioLegend) or Alexa Fluor 568 isocyanate IB4 (Life Technologies) for blood vessel visualization, Alexa Fluor 647 anti-CD68 (AbD Serotec) for monocyte/macrophage visualization, and Alexa Fluor 488 anti-CD206 (AbD Serotec) or Alexa Fluor 647 anti-CD206 (Biolegend). Tissues were mounted in 50/50 glycerol/phosphate buffered saline and imaged through the entire thickness of the muscle (~200 microns) on a Zeiss LSM 710 NLO confocal microscope or a Nikon confocal microscope.

For monocyte/macrophage quantification, 3–4 different fields of view (FOVs) per muscle containing a collateral arteriole with monocytes/macrophages evident were located manually. Full-thickness z-stack (2 μm step size) volume renders of these FOVs were generated using a 20X oil immersion objective. 20X confocal z-stack volume renders were used for cell association quantification. A threshold at which only the brightest CX3CR1<sup>+</sup> cells (CX3CR1<sup>hi</sup>) were visible was applied to all images. Vessel-associated CD206<sup>+</sup> and CX3CR1<sup>lo</sup> cells were defined as cells falling within the 2-dimensional area of the vessel in question and within 50 μm of the vessel border. ImageJ (NIH) imaging software was used to quantify vessel length and density. Arterioles were distinguished from venules by degree of lectin binding, vessel size, and morphology. Cell counts and green channel thresholding to designate CX3CR1<sup>hi</sup> cells were performed in Adobe Photoshop (Adobe Systems Incorporated).

Angiogenesis assay with monocyte co-culture. C166 murine yolk-sac endothelial cells (ATCC) were propagated in flasks coated with 0.1% gelatin (Stem Cell Technologies) using endothelial growth medium (Angioproteomie) and incubated at 37 °C and 5% CO<sub>2</sub> atmosphere. Monocytes were isolated from bone marrow of male CX3CR1<sup>GFP<sup>+</sup>/</sup> mice and sorted using a FACS-Aria IIIu to discriminate CX3CR1<sup>hi</sup> monocytes and CX3CR1<sup>lo</sup> monocytes. Endothelial tube forming assays were performed in 15-well angiogenesis slides (Ibidi) as follows. Wells were coated with BD Matrigel<sup>TM</sup> and seeded with C166 cells (7500 cells/well) and endothelial cells. After 24 hours of cell incubation, monocytes were added to the C166 cell–endothelial cell co-culture and imaged through the entire thickness of the muscle (~200 microns) on a Zeiss LSM 710 NLO confocal microscope.

Statistical analysis. Data are presented as mean ± standard error of the mean (S.E.M.). All statistical analysis was performed in GraphPad Prism software. Comparisons of two groups were made using a two-tailed unpaired t-test, with Welch's correction if standard deviations were not equal. For studies with two independent variables, two-way ANOVA with Sidak's test for multiple comparisons was performed. For confocal intravital microscopy (Fig. 6), data reflect eight ROIs acquired across 3–4 animals per group, and statistical comparisons were made using a two-tailed Mann-Whitney test. Unless otherwise noted, p < 0.05 was considered statistically significant.

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Author Contributions
C.E.O., C.L.S.E., M.E.O., and J.R.K. designed and performed in vivo studies, analyzed the data, and wrote the manuscript. A.C.B. performed experiments, analyzed data, and provided critical feedback. D.P.P. performed in vitro assays, designed computer simulations, analyzed data, and reviewed the manuscript. B.T.J. performed data analysis and reviewed the manuscript. E.A.B. and S.M.P. provided resources, designed experiments, wrote the manuscript, and supervised the project.

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