Hematopoietic stem cells (HSCs) remain mostly quiescent under steady-state conditions but switch to a proliferative state following hematopoietic stress, e.g., bone marrow (BM) injury, transplantation, or systemic infection and inflammation. The homeostatic balance between quiescence, self-renewal, and differentiation of HSCs is strongly dependent on their interactions with cells that constitute a specialized microanatomical environment in the BM known as the HSC niche. Here, we identified the secreted extracellular matrix protein Del-1 as a component and regulator of the HSC niche. Specifically, we found that Del-1 was expressed by several cellular components of the HSC niche, including arteriolar endothelial cells, CXCL12-abundant reticular (CAR) cells, and cells of the osteoblastic lineage. Del-1 promoted critical functions of the HSC niche, as it regulated long-term HSC (LT-HSC) proliferation and differentiation toward the myeloid lineage. Del-1 deficiency in mice resulted in reduced LT-HSC proliferation and infringed preferentially upon myelopoiesis under both steady-state and stressful conditions, such as hematopoietic cell transplantation and G-CSF– or inflammation-induced stress myelopoiesis. Del-1–induced HSC proliferation and myeloid lineage commitment were mediated by β3 integrin on hematopoietic progenitors. This hitherto unknown Del-1 function in the HSC niche represents a juxtacrine homeostatic adaptation of the hematopoietic system in stress myelopoiesis.
Secreted protein Del-1 regulates myelopoiesis in the hematopoietic stem cell niche

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Hematopoietic stem cells (HSCs) remain mostly quiescent under steady-state conditions but switch to a proliferative state following hematopoietic stress, e.g., bone marrow (BM) injury, transplantation, or systemic infection and inflammation. The homeostatic balance between quiescence, self-renewal, and differentiation of HSCs is strongly dependent on their interactions with cells that constitute a specialized microanatomical environment in the BM known as the HSC niche. Here, we identified the secreted extracellular matrix protein Del-1 as a component and regulator of the HSC niche. Specifically, we found that Del-1 was expressed by several cellular components of the HSC niche, including arteriolar endothelial cells, CXCL12-abundant reticular (CAR) cells, and cells of the osteoblastic lineage. Del-1 promoted critical functions of the HSC niche, as it regulated long-term HSC (LT-HSC) proliferation and differentiation toward the myeloid lineage. Del-1 deficiency in mice resulted in reduced LT-HSC proliferation and infringed preferentially upon myelopoiesis under both steady-state and stressful conditions, such as hematopoietic cell transplantation and G-CSF– or inflammation-induced stress myelopoiesis. Del-1–induced HSC proliferation and myeloid lineage commitment were mediated by β3 integrin on hematopoietic progenitors. This hitherto unknown Del-1 function in the HSC niche represents a juxtacrine homeostatic adaptation of the hematopoietic system in stress myelopoiesis.

Introduction

Hematopoietic stem cells (HSCs) lie at the core of the hematopoietic and immune systems and have a unique capacity for self-renewal and differentiation to multipotent and lineage-committed hematopoietic progenitors, which give rise to mature cells of the myeloid and lymphoid lineages (1). Under steady-state conditions, HSCs remain mostly quiescent and have a limited function in the maintenance of hematopoiesis (1–3). However, HSCs promptly switch to a proliferative state in order to meet the increased demands in response to hematopoietic stress, such as bone marrow (BM) injury, transplantation, or systemic infection (1, 3). The specialized micro-anatomical environment in the BM in which HSCs reside is designated the HSC niche and crucially regulates the homeostatic balance between quiescence, self-renewal, and differentiation of HSCs. Recent advances in BM imaging reveal a perivascular distribution of HSCs in their niche, which comprises several cell populations. Endothelial (4–11) and perivascular stromal cells (9, 12–14), including CXCL12-abundant reticular cells (CAR cells) (15, 16), are considered the main cell populations constituting the homeostatic HSC niche, whereas osteolineage cells and osteoblasts have a significant role in the post-transplant HSC niche (3, 17, 18). Niche cells regulate HSC homeostasis through paracrine growth factor and chemokine signals as well as via adhesive interactions (3, 5, 9, 19, 20).

Therapies targeting HSCs, including the administration of growth factors such as granulocyte colony–stimulating factor (G-CSF) and HSC transplantation, have revolutionized the clinical outcome in patients with malignancies or BM failure syndromes (21, 22). G-CSF–based HSC mobilization is the main strategy used in both autologous and allogeneic transplantation (23). However, G-CSF administration fails to mobilize adequate numbers of HSCs for transplantation in a substantial number of healthy donors and patients (21, 23, 24), rendering the development of additional mobilization strategies imperative. Moreover, G-CSF is used as a prophylactic treatment in patients with increased risk for iatrogenic neutropenia (25, 26) and in the treatment of patients with congenital neutropenia (27, 28).

Developmental endothelial locus–1 (Del-1), a glycoprotein secreted by endothelial and certain other cells, becomes associated with the extracellular matrix or cellular surfaces and interacts with
We show that Del-1, via its interaction with the αvβ3 integrin, promotes several critical functions in the niche, including HSC retention, hematopoietic progenitor cell cycle progression, and myeloid lineage commitment of HSCs. Del-1 thereby regulates myelopoiesis under steady-state conditions and in G-CSF– or inflammation-induced stress myelopoiesis, as well as myelopoiesis reconstitution under regenerative/transplantation conditions. Del-1 is hence a niche component that serves a juxtacrine homeostatic adaptation of the hematopoietic system in inflammation-related and regeneration myelopoiesis.

Results
Del-1 expression in the BM. First, we sought to investigate whether Del-1 is present in the BM. We initially studied the expression of the Del-1–encoding gene Edil3 in the BM niche and hematopoietic cell populations. We found that Edil3 mRNA expression was significantly higher in the endosteal region as compared with the central BM (cBM) (Figure 1A), suggesting that Del-1 is enriched at the endosteal area of the BM. Analysis of sorted cells from CXCL12-GFP mice demonstrated that Edil3 was highly expressed in stromal cell populations from CXCL12-GFP mice: CD45–Ter119–CD31–GFPhi (CAR) cells, CD45–Ter119–CD31–MSCs, and CD45–Ter119–CD31–GFP+ endothelial cells (EC; n = 3–4). The mRNA expression was normalized against β2M. (C) Gating strategy for the isolation of endothelial cells. After gating on CD45–Lin– cells, sinusoidal (sBMEC; CD31–Sca1+) and arteriolar (aBMEC; CD31–Sca1+) BM endothelial cells were isolated. VE-cadherin (VE-cadh) staining was used to confirm the arteriolar origin of the CD31–Sca1+ cell population. Right: Edil3 mRNA levels in sBMECs and aBMECs (n = 3–4). mRNA expression was normalized against β2M. (D) aBMECs were further sorted according to Vcam1 expression. Edil3 mRNA levels in Vcam1lo and Vcam1hi aBMECs as well as in CD45–Lin–CD31–Sca1–CD51+ OSL cells (n = 4–5). mRNA expression was normalized against β2M. (E) Localization of Del-1 in the perivascular area of the BM; vessel lumen staining was performed with isolectin B4 (lectin). Del-1–deficient mice served as controls for the Del-1 staining. (F) Fluorescence microscopy image showing the presence of Del-1 in arterioles. Endothelial staining was performed using anti-PECAM1 and VE-cadherin antibodies. Scale bars: 5 μm. (G) Edil3 mRNA in hMSCs and primary human osteoblasts (hOBs) was compared with Edil3 mRNA in HUVECs (HUVECs, n = 4 independent cultures; hMSCs, n = 4 donors; hOBs, n = 1 performed in technical replicates). The mRNA expression was normalized against GAPDH. (H) Del-1 concentration in culture supernatants of hMSCs was assessed by ELISA (n = 4 donors). Data are presented as mean ± SEM. Mann-Whitney U test, *P < 0.05, **P < 0.01.
in CAR cells (CD45^TER119 CD31^GFP^hi) as compared with endothelial cells (CD45^TER119 CD31^GFP^lo) and CXCL12-negative stromal cells (aBMECs) as compared with Lin^−CD45^−CD31^−Sca1^−CD51^hi osteolineage (OSL) cells (Figure 1D). In contrast, Edil3 mRNA expression could not be detected in multipotent progenitors (MPPs; Lin^−cKit^Sca1^−CD48^−CD150^), long-term HSCs (LT-HSCs) (Lin^−cKit^Sca1^−CD48^−CD150^), or other hematopoietic cell populations in the BM, such as Lin^−^ cells (data not shown). The spatial distribution of Del-1 in the perivascular area of the BM and its expression in the endothelium of BM arterioles (diameter approximately 5 µm) were verified by immunostaining (Figure 1E, F, and F, and Supplemental Figure 1). We further evaluated the expression of Del-1 in primary human BM-derived mesenchymal stromal cells (hMSCs) and primary human osteoblasts (hOBs) by quantitative PCR. 

Del-1 expression was highly expressed in hMSCs and hOBs, compared with HUVECs (Figure 1G), a cell population known to express substantial levels of Del-1. This expression was confirmed by RT-qPCR in CAR cells (CD45^TER119 CD31^GFP^lo) and CXCL12-negative stromal cells, like CAR cells, (9, 14–16), and by cells that mediate the reconstitution of hematopoiesis after transplantation, like OSL cells (3, 17, 18).

**Del-1 promotes steady-state myelopoiesis.** We next assessed whether Del-1 could affect hematopoietic progenitor maintenance and function. Specifically, to determine a possible functional role of Del-1 in the regulation of hematopoiesis, we performed BM analysis in adult Del-1-deficient (Del^−/−^) and Del-1-proficient (WT; Del^+/+^) mice. Although Del-1 deficiency had no significant effect on total BM cellularity (Figure 2A), it caused a significant decrease in the number of common myeloid progenitors (CMPs; Lin^−cKit^Sca1^CD16/32^CD34^) (Figure 2B) and megakaryocyte erythroid progenitors (MEPs; Lin^−cKit^Sca1^CD16/32^CD34^) (Figure 2C). The numbers of granulocyte macrophage progenitors (GMPs; Lin^−cKit^Sca1^CD16/32^CD34^) were not significantly decreased in the BM of Del^−/−^ mice as compared with Del^+/+^ mice (Figure 2D). Moreover, the use of a CFU assay revealed decreased numbers of functional progenitor cells (colony-forming cells [CFCs]) in the BM of Del^−/−^ mice, as compared with Del^+/+^ mice (Figure 2E). The decrease in myeloid progenitor cells was associated with a significant decrease in the numbers of Gr1^hi^CD11b^+ granulocytes (Figure 2, F and G) and Gr1^int^CD11b^+ myeloid cells (Figure 2, F and H) in the BM of Del^−/−^ mice as compared with Del^+/+^ mice. In contrast, no difference was observed in the levels of G-CSF in peripheral blood (Supplemental Figure 2), in the number of different B cell populations, or in the numbers of common lymphoid progenitors (CLPs; Lin^−Sca1^cKit^Flt3^IL7Ra^) in the BM due to Del-1 deficiency (Supplemental Figure 3). Since steady-state hematopoiesis is maintained by long-lived multipotent progenitors (2), we studied whether Del-1 deficiency had a quantitative and/or a qualitative effect on hematopoietic progenitors. No difference was seen in the
Together, these data suggest that Del-1 promotes steady-state myelopoiesis by regulating the bias of early hematopoietic progenitors toward the myeloid lineage.

To assess whether the decreased numbers of myeloid progenitor cells in the BM of Del-1–deficient mice were due to impaired retention of hematopoietic and myeloid progenitor cells in the BM, we analyzed LSK cells and myeloid progenitor cells in the spleen, as well as circulating LSK cells, myeloid progenitors (LK; Lin–Sca1–cKit+), and CFCs. We did not observe any difference in circulating progenitors (Supplemental Figure 5, A–C), whereas the numbers and frequency of both LSK and LK cells were decreased in the spleen of Del-1–deficient mice (Supplemental Figure 5, D–G). These data exclude the possibility that the decrease in myeloid progenitor cells in the Del-1–deficient BM was due to enhanced mobilization at steady state and, moreover, suggest a role for Del-1 in promoting HSC differentiation toward the myeloid lineage.

Del-1 promotes the regeneration of myelopoiesis after hematopoietic cell transplantation. We next studied whether Del-1 deficiency
could affect the long-term lineage output of transplanted hematopoietic progenitors. To this end, we generated BM chimeras by transplanting CD45.1+ BM cells from congenic B6.SJL (CD45.1+) mice into lethally irradiated Edil3+/+ or Edil3−/− mice (CD45.2) (Figure 3A). At 16 weeks after transplantation, we observed a significant decrease in the percentage of donor-derived Gr1+CD11b+ myeloid cells in the peripheral blood of Edil3−/− mice compared with Edil3+/+ recipient mice (Figure 3B), with a corresponding increase in the percentage of B cells (Figure 3C). Analysis of BM revealed that this difference was due to impaired reconstitution of myeloipoiesis in Edil3−/− compared with Edil3+/+ recipients (Figure 3, D–F), whereas the regeneration of the lymphoid lineage in the BM remained unaffected (Figure 3G). To definitively determine whether Del-1 can affect the long-term lineage bias of transplanted hematopoietic progenitors, we re-transplanted donor-derived cells (CD45.1+) from the Edil3+/+ or Edil3−/− mice into secondary Edil3+/+ and Edil3−/− recipients. The observed impairment in myeloid lineage reconstitution in the peripheral blood of secondary Edil3−/− recipients was strikingly exacerbated (Figure 3, H and I), indicating that Del-1 in the BM drives the differentiation
of hematopoietic progenitors, particularly LT-HSCs, toward the myeloid lineage.

After transplantation, hematopoietic progenitor cells localize in proximity to blood vessels at the endosteal area of the BM (3, 17, 18, 39), which could be attributed to irradiation-induced disruption of sinusoids, though not of endosteal arterioles (3). The expression of Del-1 in the peri-arteriolar region and by OSL cells, which also contribute to the restoration of hematopoiesis after transplantation, prompted us to further investigate in more detail how Del-1 supports the regeneration of myelopoiesis. To this end, we generated BM chimeras by transplanting CD45+ BM cells from congenic B6.SJL (CD45.1+) mice into lethally irradiated Edil3–/– or Edil3+/+ mice (CD45.2+) as above and performed BM analysis at 6 weeks after transplantation. Del-1 deficiency in the BM impaired the expansion of donor-derived hematopoietic cells, as shown by the decreased numbers of donor-derived LSK cells (Figure 4, A and B), ST-HSCs (Figure 4, C and E), and MPPs (Figure 4, C and F), whereas LT-HSC numbers were not altered (Figure 4D). Del-1 deficiency in recipients was associated with an increased percentage of donor-derived LT-HSCs and a concomitant decrease in the percentage of MPPs (Figure 4G), thus pointing to potentially diminished differentiation-associated proliferation of HSCs in the Del-1-deficient recipient environment. These findings were further supported by a decrease in the numbers of myeloid progenitor cells, CMPs and GMPs, in Edil3–/– as compared with Edil3+/+ recipient mice (Figure 4, H–J). These data together suggest that Del-1 promotes the differentiation of hematopoietic progenitor cells into the myeloid lineage after transplantation. Additionally, Del-1 deficiency resulted in reduced numbers of Gr1hiCD11b+ granulocytes (Figure 4K) and Gr1intCD11b+ myeloid cells (Figure 4L), whereas B cell numbers were not affected (Figure 4M). Upon transplantation, therefore, Del-1 promotes hematopoietic progenitor engraftment and differentiation toward the myeloid lineage.

Del-1 deficiency increases LT-HSC quiescence. Since normally there is a strong imbalance in the differentiation output of hematopoietic progenitor cells toward myelopoiesis compared with lymphopoiesis (2, 40), we next investigated whether Del-1 deficiency could affect the proliferation potential of hematopoietic progenitors and thereby the progression of differentiation toward myeloid cell lineage. Cell cycle analysis revealed an increase in LT-HSC
bers of donor BM cells derived from either Edil3–/– competitive transplantation assay by transplanting equal number of donor BM cells from either Edil3–/– or Edil3+/+ mice. (B) Representative flow cytometry plots of donor-derived chimera in the peripheral blood of recipient mice at week 16 after transplantation and (C) number of R.U. per femur of Edil3–/– or Edil3+/+ donor mice (n = 10 recipients per group). (D) Representative flow cytometry plots and (E) percentage of donor-derived (CD45.2+) Gr1+CD11b+ myeloid cells in the blood of recipient mice. (F) Frequency of donor-derived cells in the BM of recipient mice at 16 weeks after transplantation (n = 10 recipients per group). (G) Representative flow cytometry plots and (H) percentage of donor-derived (CD45.2+) Gr1+CD11b+ myeloid cells in the BM of recipient mice (n = 10 recipients per group). (I) Representative flow cytometry plots and (J) percentage of donor-derived CD19+ B cells in the BM of recipient mice at 16 weeks after transplantation (n = 10 recipients per group). Data are presented as mean ± SEM. Mann-Whitney U test, *P < 0.05, **P < 0.01, ***P < 0.001.

Figure 6. Del-1 deficiency enhances long-term reconstitution capacity and lineage bias of LT-HSCs. (A) Competitive repopulation assay: BM cells from either Edil3–/– or Edil3+/+ mice (CD45.2+) or CD45.1+ BM cells as competitors were transplanted into B6.SJL recipient mice (CD45.1). (B) Representative flow cytometry plots of donor-derived chimera in the peripheral blood of recipient mice at week 16 after transplantation and (C) number of R.U. per femur of Edil3–/– or Edil3+/+ donor mice (n = 10 recipients per group). (D) Representative flow cytometry plots and (E) percentage of donor-derived (CD45.2+) Gr1+CD11b+ myeloid cells in the blood of recipient mice. (F) Frequency of donor-derived cells in the BM of recipient mice at 16 weeks after transplantation (n = 10 recipients per group). (G) Representative flow cytometry plots and (H) percentage of donor-derived (CD45.2+) Gr1+CD11b+ myeloid cells in the BM of recipient mice (n = 10 recipients per group). (I) Representative flow cytometry plots and (J) percentage of donor-derived CD19+ B cells in the BM of recipient mice at 16 weeks after transplantation (n = 10 recipients per group). Data are presented as mean ± SEM. Mann-Whitney U test, *P < 0.05, **P < 0.01, ***P < 0.001.

quiescence due to Del-1 deficiency, as shown by the increased percentage of LT-HSCs in G0 phase and the decreased percentage of LT-HSCs in the S/G1/M fraction in Edil3–/– mice (Figure 5, A and B). A BrdU incorporation assay further confirmed the decreased proportion of proliferating LT-HSCs in Edil3–/– mice (Figure 5, C and D). The increased quiescence of LT-HSCs from Edil3–/– mice compared with LT-HSCs from Edil3+/+ mice was associated with a downregulation in the gene expression of cyclins Ccng2, Ccnd1, and Ccng1 in Edil3–/– compared with Edil3+/+ mice (Figure 5E). Moreover, using a BrdU pulse-chase assay that can identify dormant LT-HSCs (41) (Figure 5F), we found increased frequency of BrdU-retaining (hence dormant) LT-HSCs on day 70 after BrdU withdrawal in Edil3–/– as compared with Edil3+/+ mice (Figure 5G), consistent with the increased quiescence of LT-HSCs in Del-1 deficiency.

Since the engrafment potential of hematopoietic progenitor cells mainly relies on the fraction of cells in the G0 phase of cell cycle (42, 43), we further assessed whether the enhanced quiescence of LT-HSCs from Edil3–/– mice could result in the increased engrafment capacity of these cells. To this end, we performed a competitive transplantation assay by transplanting equal number of donor BM cells derived from either Edil3–/– or Edil3+/+ mice (CD45.2+) and competitor cells (CD45.1+) into lethally irradiated recipients (CD45.1) (Figure 6A). Peripheral blood analysis at 16 weeks after transplantation revealed a significantly increased number of reconstitution units (R.U.) in the BM of Edil3–/– donors as compared with their Edil3+/+ counterparts (Figure 6, B and C). We further assessed the lineage bias of donor cells and observed a decreased frequency of donor-derived (CD45.2+) Gr1+CD11b+ myeloid cells in the blood of recipients transplanted with cells from Edil3–/– mice (Figure 6, D and B). Analysis of the BM chimerism in recipient mice further confirmed the increased long-term reconstitution efficiency of progenitors derived from Edil3–/– donors (Figure 6F). Additionally, we observed a decreased frequency of Gr1+CD11b+ myeloid cells (Figure 6, G and H) and a corresponding increase in the frequency of CD19+ B cell (Figure 6, I and J) in the donor-derived cell population in the BM of recipients transplanted with cells from Edil3–/– mice. Thus, the increased quiescence of LT-HSCs due to Del-1 deficiency results in enhanced repopulation activity of these cells and is associated with decreased myeloid differentiation potential of long-term repopulating cells.

β3 Integrin mediates Del-1–dependent LT-HSC proliferation and myeloid differentiation. Having demonstrated that Del-1 promotes cell cycle progression and myeloid differentiation of hematopoietic progenitors, we set out to address the underlying mechanisms. Through cell-cell and cell-matrix interactions, integrins act as mechanosensors that initiate cell cycle progression via regulation of the levels of cyclins and cyclin-dependent kinase inhibitors (44, 45). In previous studies, we identified the integrins αLβ2 (CD11a/CD18) and αvβ3 (CD51/CD61) as Del-1 receptors (29–31). Therefore, we employed an adhesion assay to determine whether any of these integrins on hematopoietic progenitors mediates interaction with Del-1. Blocking the β3 integrin subunit with anti-CD61 antibody significantly diminished LSK cell adhesion to immobilized Del-1, whereas blockade of CD11a or CD49d had no effect (Figure 7A). We therefore next analyzed CD61 expression on hematopoietic progenitors with flow cytometry and found that CD61 was highly expressed on LT-HSCs and to a lesser extent on MPPs (Figure 7B and C), which is in accordance with earlier studies (46, 47). To test the hypothesis that the interaction of Del-1 with the αvβ3 integrin on hematopoietic progenitors regulates cell cycle–related gene expression, we cultured isolated LSK cells on immobilized Del-1 expressed as an Fc fusion protein (Del-1–Fc) or Del-1[RGE]–Fc, a point mutant that precludes interaction with αvβ3 integrin due to a Glu (E) for Asp (D) substitution in the critical
RGD motif. Fc protein was used as a negative control. In contrast to the WT molecule, the RGE point mutant failed to induce Ccnd1 gene expression in LSK cells (Figure 7, D and E), thus indicating a requirement for αβ3-dependent adhesion to Del-1 for induction of Ccnd1. To independently confirm that signaling induced through Del-1 ligation of αβ3 integrin was responsible for regulating cyclin D1 expression, we used an inhibitor of integrin-linked kinase (ILK), which mediates integrin-dependent outside-in sig-
it had no effect on the differentiation of MPPs or CMPs toward GMPs (Supplemental Figure 6). These experiments firmly establish that the direct interaction of LT-HSC with Del-1 instructs their differentiation toward the myeloid lineage.

We next addressed the in vivo relevance of the Del-1/β3 integrin interaction. To this end, we initially studied the role of Del-1 in hematopoietic progenitor cell homing to the BM. Fluorescently labeled Lin− cells were transplanted to non-ablated Edil3−/− and Edil3+/+ mice, and the percentage of transferred LSK cells in the BM was assessed 3 or 20 hours later. Del-1 deficiency had no effect at the early time point (Supplemental Figure 7). Therefore, Del-1 does not affect the initial recruitment of hematopoietic progenitors via the vasculature to the BM, consistent with the fact that Del-1 does not interact with or regulate the α4β1 integrin (29) (Figure 7A), which is required for this function (48). Intriguingly, at the 20-hour time point, the percentage of transferred cells was significantly decreased in Edil3−/− as compared with Edil3+/+ mice, hence implicating Del-1 in the homing of transplanted hematopoietic progenitors (Figure 7M). To assess the significance of β3 integrin-dependent interactions of Del-1 in the BM, we tested whether this

Figure 8. Del-1 mediates the restoration of myelopoiesis in the BM upon LPS-induced inflammation. (A) Gr1hiCD11b+ granulocytes and (B) Gr1intCD11b+ myeloid cell numbers in the BM of Edil3−/− or Edil3+/+ mice at 24 hours (gray background) and at 72 hours following the second injection of LPS (n = 4–10 mice per group). (C) CMP, (D) LSK, (E) LT-HSC, (F) ST-HSC, and (G) MPP cell numbers in the BM of Edil3−/− and Edil3+/+ mice (n = 4–8 mice per group) at 24 hours (gray background) and at 72 hours following the second injection of LPS. (H) Cell cycle analysis in LT-HSCs and (I) MPPs at 72 hours following the second injection of LPS (n = 5–7 mice per group). Data presented as mean ± SEM. Mann-Whitney U test. *P < 0.05.

naling. We found that ILK inhibition abrogated the effect of the αvβ3–Del-1 interaction on Cend1 expression (Figure 7E). These findings reveal a role for Del-1 in regulating cyclin D1 expression in progenitors via interaction with αvβ3 integrin and integrin-dependent outside-in signaling events.

To further study whether Del-1 can drive the proliferation and differentiation of early progenitor cells, we performed an in vitro differentiation assay. LT-HSCs were treated with Fc-control, Del-1-Fc, or Del-1[RGE]–Fc in cell suspension cultures (Figure 7F). Del-1-Fc (but not Fc-control) induced the proliferation and differentiation of LT-HSCs toward GMPs, whereas this effect was abolished when the RGE point mutant was used (Figure 7G–I). Inhibition of β3 integrins by a β3 inhibitor, cileg tide, also abrogated LT-HSC differentiation on day 7 (Figure 7I), further suggesting the involvement of β3 integrins in Del-1–dependent induction of LT-HSC differentiation. To further determine whether Del-1 acts on LT-HSCs or at later stages of differentiation, we treated LT-HSCs, MPPs, or CMPs for 48 hours with Fc-control or Del-1–Fc (Figure 7K and Supplemental Figure 6). We observed that Del-1 induced the differentiation toward MPPs (Figure 7L), whereas it had no effect on the differentiation of MPPs or CMPs toward GMPs (Supplemental Figure 6). These experiments firmly establish that the direct interaction of LT-HSC with Del-1 instructs their differentiation toward the myeloid lineage.

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Integrin αvβ3 has been previously shown to mediate the maintenance of HSCs, although the natural ligand(s) required for this niche interaction were not identified (46, 47). Having shown that the Del-1/β3 integrin interaction plays a role in the homing of hematopoietic progenitors into the BM, we then determined whether this interaction also contributes to the restoration of myelopoiesis after transplantation. For this purpose, CD45+ BM cells from Itgb3−/− mice were transferred into lethally irradiated Edil3−/− or Edil3+/+ mice, and BM analysis in the recipient mice was performed 6 weeks later (Supplemental Figure 8A). As this experiment was not done using the congenic CD45.1/CD45.2 system, it was not possible to specifically study the reconstitution by donor-derived cells. For this reason, analysis was performed by determining the absolute cell numbers of different hematopoietic progenitor populations in the BM of recipient mice. Consistent with the above-described effects of β3 integrin inhibition, in the absence of β3 integrin on donor cells, Del-1 deficiency in recipients failed to affect the number of LSK cells, myeloid progenitors, or Gr1 hiCD11b+ granulocytes and Gr1 intCD11b+ myeloid cell populations (Supplemental Figure 8, B–F). These findings stand in stark contrast to the negative effect of Del-1 deficiency on myelopoiesis reconstitution upon transplantation of β3 integrin–sufficient donor cells (Figure 4). Therefore, β3 integrin is required for the Del-1–mediated progenitor engraftment and reconstitution of myelopoiesis after transplantation.

Del-1 promotes the response of hematopoietic progenitors to systemic inflammation. The above-documented function of Del-1 in steady-state myelopoiesis and in the regeneration of the myeloid lineage after transplantation prompted us to investigate its possible involvement in the response of hematopoietic progenitors to stress granulopoiesis, such as in models induced by LPS or G-CSF.

Administration of high-dose LPS is widely used to mimic the effect of systemic microbial infection on granulopoiesis, which forces LSK proliferation to compensate for the increased need for mature myeloid cells and restore BM cellularity (49–51). We therefore engaged this model to investigate whether Del-1 in the niche regulates the potential for myeloid lineage replenishment in the BM. To this end, Edil3−/− or Edil3+/+ mice were intraperito-

**Figure 9. Role of Del-1 in the response of hematopoietic progenitor cells to G-CSF administration.**

(A) Total cell numbers and (B) LSK, (C) LT-HSC, (D) ST-HSC, (E) MPP, (F) CMP, and (G) GMP cell numbers in the BM of Edil3−/− and Edil3+/+ mice (n = 6 per group) at the end of the G-CSF administration protocol. (H) CFCs from the BM of Edil3−/− (n = 6) and Edil3+/+ mice (n = 5) at the end of the G-CSF administration protocol. (I) Representative flow cytometry plots and (J) numbers of mobilized LSK cells in the peripheral blood of Edil3−/− (n = 10) and Edil3+/+ mice (n = 11) at the end of the G-CSF administration protocol. (K) Number of CFCs in the peripheral blood of Edil3−/− and Edil3+/+ mice (n = 5-6 mice) at the end of the G-CSF administration protocol. Data are presented as mean ± SEM. Mann-Whitney U test, *P < 0.05, **P < 0.01.
neally injected with ultra-pure LPS twice, with an interval of 48 hours, and hematopoietic progenitors and myeloid lineage cells were analyzed at 24 hours and at 72 hours following the second injection (52, 53). In this model, the initial increase in peripheral blood myeloid cells (52, 53), associated with a decrease in the numbers of mature myeloid cells and CMPs in the BM, is followed by a prominent expansion of the myeloid cell compartment, as observed in Edil3–/– mice at 72 hours after LPS administration (Figure 8B). However, the restoration of myelopoiesis in the BM was significantly attenuated in Del-1 deficiency, as shown by the decreased numbers of Gr1hiCD11b+ granulocytes (Figure 8A) and Gr1hiCD11b+ myeloid cells (Figure 8B) in Edil3–/– relative to Edil3+/+ mice. In the same vein, the numbers of CMPs were decreased in Edil3+/+ compared with Edil3–/– mice at 72 hours after LPS administration (Figure 8C). Furthermore, Del-1 deficiency resulted in relatively inefficient expansion of LSK cells caused predominantly by inefficient expansion of MPPs 24 hours after LPS injection, and these differences between Edil3–/– and Edil3+/+ mice were sustained after 72 hours (Figure 8, D–G). The delay in the reconstitution of the myeloid lineage was also associated with an increase in the percentage of quiescent LT-HSCs and MPPs (Figure 8, H and I) after 72 hours. In contrast, no difference was observed in cell cycle progression of myeloid progenitor cells upon LPS injection due to Del-1 deficiency (data not shown). No alterations in G-CSF production, which plays a critical role in emergency granulopoiesis (54), were observed between Edil3–/– and Edil3+/+ mice (Supplemental Figure 9). In addition, the expression of Edil3 in the BM was not altered after LPS administration (data not shown). Furthermore, we did not observe any difference in the number of LSK and LK cells in the peripheral blood or spleen between Del-1–deficient and –proficient mice (Supplemental Figure 10). The data in toto demonstrate that Del-1 promotes myeloid cell replenishment in the BM in the course of systemic inflammation.

Del-1 regulates hematopoietic progenitor proliferation and retention in response to G-CSF. Besides inducing myeloid progenitor maturation (54), G-CSF was previously shown to promote LSK cell expansion by affecting the LT-HSC cell cycle (55, 56). For this reason, adult Edil3–/– and Edil3+/+ mice were injected daily with G-CSF for up to 6 days. At 24 hours after a single G-CSF administration, the expansion of total BM cells (Supplemental Figure 11A), Gr1hiCD11b+ granulocytes (Supplemental Figure 11B), and Gr1hiCD11b+ myeloid cells (Supplemental Figure 11C) was less prominent in Edil3–/– than in Edil3+/+ mice. LSK cell, MPP, and CMP but not LT-HSC or ST-HSC numbers (Supplemental Figure 11, D–H) were also decreased in Edil3–/– relative to Edil3+/+ mice. Cell cycle analysis of LT-HSCs from Edil3–/– and Edil3+/+ mice after 3 daily injections of G-CSF revealed a decrease in the proliferation potential in HSCs derived from Edil3–/– mice (Supplemental Figure 11, I and J). By the end of the G-CSF administration course (6 days), BM cellularity (Figure 9A) and numbers of hematopoietic and myeloid progenitors (Figure 9, B–G) were dramatically reduced in Edil3–/– mice as compared with Edil3+/+ mice. The decrease in phenotypic progenitor cells in Del-1 deficiency by the end of the G-CSF administration course was also associated with reduced counts of functional progenitor cells, as shown by a CFU assay (Figure 9H). To further evaluate whether Del-1 deficiency affects myelopoiesis upon G-CSF administration, the percentage of GMPs and LSK cells in the BM after 6 daily injections of G-CSF was expressed relative to the percentage of GMPs and LSK cells in the BM after 6 daily injections of PBS. The corresponding ratios were then compared between Edil3–/– and Edil3+/+ mice. We observed that the expansion of GMPs in the BM of Edil3–/– mice was more robust compared with Edil3+/+ mice (Supplemental Figure 12A). Additionally, the relative levels of LSK cells were also significantly higher in Edil3–/– mice (Supplemental Figure 12B).

As G-CSF is used as a mobilizing agent in the context of autologous or allogeneic transplant (21, 23), we examined whether the physical interaction between Del-1 and HSCs promotes their retention in the BM, thereby counteracting the mobilizing effect of G-CSF. G-CSF–dependent mobilization of hematopoietic progenitor cells was more efficient in Del-1 deficiency, as shown by the increased numbers of LSK (Figure 9, I and J) and CFUs (Figure 9K) in the peripheral blood of Edil3–/– mice. We did not observe any alterations in the expression of Edil3 in the endosteal region of mice by the end of the G-CSF administration protocol (data not shown). Since downregulation in the expression of HSC niche factors in MSCs and osteoblasts is an important event during G-CSF–dependent mobilization (12, 57, 58), we assessed Cxcl12, Angpt1, and Kitl gene expression in the endosteal region of Edil3–/– and Edil3+/+ mice after 6 daily injections of G-CSF. We observed that the decrease in the mRNA levels of these factors followed the same pattern in Edil3–/– and Edil3+/+ mice (Supplemental Figure 13). Therefore, the effect of Del-1 deficiency on G-CSF-induced mobilization cannot be attributed to alterations in the expression of other HSC niche factors. These findings overall indicate that Del-1 acts as a niche factor that supports hematopoietic progenitor retention in the BM upon G-CSF–induced mobilization.

Discussion

Herein, we detected that Del-1 was enriched in the perivascular area of the BM, expressed by critical cellular niche components. Moreover, Del-1, via its interaction with β3 integrin on LT-HSCs, facilitated LT-HSC proliferation and differentiation toward the myeloid lineage under steady-state and hematopoietic stress conditions. Our data therefore establish a role for Del-1 in the HSC niche microenvironment.

During steady-state hematopoiesis, Del-1 deficiency resulted in increased quiescence of LT-HSCs without affecting hematopoietic progenitor cell numbers, which is in accordance with previous studies showing that deficiency of niche factors does not necessarily have a quantitative effect on LT-HSCs (5, 59). Strikingly, however, Del-1 deficiency resulted in a significant decrease in myeloid cell populations. Steady-state myelopoiesis is mainly maintained by multipotent hematopoietic progenitor cells (2). Consistent with this finding, we detected a significant downregulation in the expression levels of myelopoiesis-related genes in CD48– LSK cells derived from Del-1–deficient mice. Moreover, Del-1 deficiency resulted in phenotypic changes in MPPs, leading to a decreased percentage of MPPs with low Fli3 expression, which have been previously shown to be biased toward differentiation into myeloid lineage (36–38). Del-1 deficiency was further shown to affect the lineage bias of LT-HSCs, as shown by competitive repopulation conditions. The enhanced engraftment potential of hematopoietic progenitors from Edil3–/– mice was linked to a diminished poten-
tial for production of myeloid cells, which is in line with recent studies showing an association between increased proliferation of LT-HSCs and myeloid differentiation bias (60, 61). In particular, HSC clones with high proliferation rates were recently shown to be biased toward myeloid differentiation, while another study revealed a correlation between the expression of cell cycle and DNA replication genes and that of myeloid differentiation genes (60, 61). Our findings that Del-1 regulated the proliferation potential and myeloid differentiation of LT-HSCs but not of MPPs or CMPs suggest that Del-1 likely promotes differentiation-associated proliferation of LT-HSCs, thereby enabling the response of hematopoiesis to increased needs for production of myeloid cells.

The effect of Del-1 deficiency on LT-HSC proliferation potential and myelopoiiesis had a strong impact in distinct models of stress myelopoiesis, wherein induction of hematopoietic progenitor proliferation is required to effectively replenish BM cells (1, 40). Using three different, clinically relevant models, we demonstrated that Del-1 is essential for the proper hematopoietic progenitor response with regard to myeloid cell reconstitution, after BM transplantation, G-CSF administration, and LPS-dependent inflammation. In all three models, Del-1 deficiency consistently compromised the ability of the niche microenvironment to support the proliferation and differentiation of LT-HSCs into the myeloid lineage. After hematopoietic cell transplantation, Del-1 in the recipient niche promoted progenitor engraftment and the generation of both progenitors and mature cells of the myeloid lineage in a manner dependent upon β3 integrin expression in hematopoietic cells. Similarly, Del-1 was found to contribute to the proliferative response of hematopoietic progenitor cells after G-CSF administration or LPS-driven inflammation. Moreover, Del-1 promoted the retention of hematopoietic progenitors in the BM upon G-CSF administration. The critical involvement of Del-1 in all these different models of stress myelopoiesis attests to the important regulatory role of Del-1 within the HSC niche for myeloid lineage commitment.

The αvβ3 integrin on LT-HSCs interacted with and mediated the Del-1-dependent progenitor retention in the BM and progenitor commitment toward the myeloid lineage. The αvβ3 integrin/Del-1 interaction supported LT-HSC differentiation-associated proliferation, which involved the induction of cyclin D1 expression. Previous studies have shown that αvβ3 integrin is expressed in HSCs and supports the maintenance of HSC activity (46, 47), although the niche-relevant ligand(s) of αvβ3 were not determined. Integrin activation is tightly linked to cell survival and proliferation (45). Cell-cell or cell–extracellular matrix adhesion through integrins induces intracellular signaling pathways that enable cells to enter cell cycle (45). The expression of D-cyclins is controlled by the extracellular environment (62) and is essential for integrin-mediated progression through the G phase of the cycle (44, 63). The important role of cyclin D1 in LT-HSC function was further highlighted in a recent study where downregulation of cyclin D1 expression, due to alterations in cell metabolism, exerted a critical impact on normal and leukemic HSC function (64). Additionally, cyclin D1 expression has been previously associated with HSC differentiation-associated proliferation (42). We demonstrate here that the decreased proliferation of LT-HSCs due to Del-1 deficiency in vivo is associated with reduced mRNA levels of Ccnd1, whereas the interaction between Del-1 and αvβ3 integrin on hematopoietic progenitors results in increased Ccnd1 expression. The Del-1 deficiency–associated decreased gene expression of Ccnd1 in hematopoietic progenitors together with the in vitro stimulatory effect of Del-1 on Ccnd1 expression strongly corroborates the evidence presented here that Del-1 acts as a niche factor mediating differentiation-associated proliferation and myeloid lineage commitment of HSCs.

Several cellular components have been implicated as contributing to homeostatic HSC niche function. Arteriolar vessels, comprising endothelial cells and a layer of mesenchymal stromal cells, promote the maintenance of LT-HSCs in a quiescent state (6, 7, 13), although non-dividing stem cells have also been localized around sinusoids (6, 10). Endothelial and perivascular stromal cells, including CAR cells, support the maintenance of HSCs in the niche by producing stem cell factor, pleiotrophin, and CXCL12 (6, 7, 9, 11-16) or by direct adhesive interactions through molecules, such as E-selectin (5). While expression of E-selectin is restricted to the endothelium, thus representing an endothelial-specific niche, Del-1 is expressed by several major cellular components in the niche microenvironment (CAR, arteriolar endothelial, and OSL cells). Therefore, Del-1 can act as a crucial HSC niche regulator in different settings. Indeed, by multiple complementary experimental approaches, we have demonstrated that Del-1 promotes the retention of hematopoietic progenitors in the niche and orchestrates their proliferation and differentiation into the myeloid lineage under diverse conditions, ranging from steady-state to regenerative and inflammation-related myelopoiesis. Thus, the demonstration of Del-1 as a factor governing the proliferation and fate of LT-HSC provides mechanistic insights into how the HSC niche operates and contributes to homeostatic adaptation of the hematopoietic system. Interestingly, the adhesive interaction between niche-expressed Del-1 and the αvβ3 integrin on LT-HSCs also constitutes an example of juxtacrine regulation of progenitor commitment toward the myeloid lineage. Our findings thereby identify what we believe to be a novel spatial component in the regulation of lineage fate and add to the growing evidence of distinct control of hematopoietic precursors within the BM.

Our findings establish a homeostatic role for Del-1 during inflammation via stimulation of myelopoiesis and production of mature effector cells. Therefore, the homeostatic function of Del-1 is not restricted to regulation of inflammatory cell recruitment to peripheral tissues as we showed earlier (29, 31, 65), but also involves control of their progenitors in the BM. As febrile neutropenia, associated with chemotherapy or HSC transplantation, is a major cause for morbidity and mortality (25), the herein identified role of Del-1 in the induction and maintenance of proper de novo granulopoiesis is important and could be therapeutically exploited in this context. On the other hand, inhibition of the interaction of Del-1 with hematopoietic progenitor cells could be exploited pharmacologically to enhance progenitor mobilization in the context of autologous or allogeneic HSC transplantation or hematologic malignancy (66). The emergence of Del-1 as a critical niche factor that promotes hematopoietic progenitor expansion and retention, as well as myelopoiesis, suggests that this homeostatic factor needs to be seriously considered in the context of BM transplantation or hematological disorders.
To generate BM chimeras, a total of $2 \times 10^6$ CD45+ BM cells from B6/SJL (CD45.1) mice were retro-orbitally transplanted into lethally irradiated (9 Gy) Edil3–/+ or Edil3+/+ (WT) littermate mice (CD45.2). Blood was collected at week 16 for analysis of donor-derived peripheral blood cells, and BM analysis was performed at 6 and 16 weeks after transplantation for the analysis of donor-derived (CD45.1) cell populations. At week 16 after transplantation, a total of $2 \times 10^6$ CD45.1 BM cells from the Edil3–/+ recipient group were further transplanted into lethally irradiated (9 Gy) Edil3–/+ or Edil3+/+ mice (CD45.2), and peripheral blood analysis of the frequency of different CD45.1+ cell populations was performed every 4 weeks. In additional experiments, $2 \times 10^6$ CD45.1 BM cells from Itgb3–/– (mouse) were retro-orbitally transplanted into lethally irradiated Edil3–/+ or Edil3+/+ mice. The reconstitution of different cell populations in the BM of the recipient mice was analyzed at week 6 after transplantation.

For BM competitive repopulation assays, the CD45.1/CD45.2 congenic system was used. Equivalent volumes of BM cells (2.5% of cells derived from a femur) harvested from CD45.2+ donor mice (Edil3–/+ and Edil3+/+) were retro-orbitally transplanted into lethally irradiated B6/SJL (CD45.1) recipients along with $3 \times 10^5$ BM cells derived from B6/SJL mice (67). Peripheral blood chimerism was assessed at 16 weeks after transplantation. We calculated the number of injected R.U. per recipient mouse using the formula ($D_j = D \times C_j / (100 - D)$), where $D$ is the number of CD45.2+ blood leukocytes at 16 weeks in the recipient mice, and $C$ is the number of competing CD45.1+ R.U. that were coinjected ($C = 3$, since $3 \times 10^5$ competing BM cells were injected). The number of R.U. per femur was calculated afterward (5, 68). Chimerism in BM cells was also analyzed at week 16. Analysis of the lineage output of donor-derived (CD45.2+) cells was further performed after gating on donor-derived cells. In all transplantation experiments, mice were kept on antibiotic-containing water for 2 weeks after irradiation.

To assess the effect of Del-1 in hematopoietic progenitor migration and homing to the BM, $2 \times 10^6$ Lin− cells from Edil3–/+ mice were labeled with CFSE and were retro-orbitally transferred into non-irradiated Edil3–/+ or Edil3+/+ littermate recipients. BM was analyzed 3 or 20 hours after cell transfer. In other experiments, CFSE+Lin− cells from Edil3–/+ mice were pretreated for 5 minutes with colchicine (10 mg/kg; Selleck Chemicals) (69) or RGD control peptide (10 mg/kg; Enzo Life Sciences, GRADSP) before their transfer into non-irradiated Edil3–/+ or Edil3+/+ littermate recipients.

Primary human cells. Primary human MSCs from healthy donors were generated out of BM aspirates by applying plastic adherence and serial passaging, as previously described (70–74). MSCs of the second passage were cultured in 24-well culture plates in DMEM supplemented with 10% FBS until reaching subconfluence, then the medium was changed to DMEM with 0.1% BSA (Sigma-Aldrich); supernatants were collected 18 and 48 hours thereafter. HUVECs (Lonza) were cultured in complete EBM-2 medium (Lonza). Cell pellets from human osteoblasts were from PromoCell. Total RNA was extracted using TRI reagent (Molecular Research Center Inc.). Del-1 concentration in culture supernatants was measured using a human EDIL3 DuoSet ELISA (R&D Systems).

**BrdU incorporation and pulse-chase assay.** For BrdU incorporation assay, BrdU was administered in drinking water for 3 days at a concentration of 0.5 mg/ml (5). For BrdU pulse-chase assay, BrdU was administered in drinking water for 15 days at a concentration of 0.5 mg/ml. Drinking water was changed twice weekly. Water containing BrdU was replaced with regular water after the pulse period, and analysis was performed 70 days after BrdU withdrawal (41). Staining for BrdU was performed using the FITC BrdU Flow Kit (BD Biosciences) following the manufacturer’s instructions.

**LPS and G-CSF administration.** To assess the effect of G-CSF on hematopoietic progenitors, we injected mice subcutaneously with a daily dose of 250 μg/kg of recombinant human G-CSF (Amen) in PBS (Life Technologies) or PBS as control for up to 6 days. For LPS-induced inflammation, mice were injected intraperitoneally with 35 μg ultrapure LPS (E. coli 0111B4; InvivoGen) twice, with a 48-hour interval, and sacrificed at 24 hours or at 72 hours following the second injection. In the LPS model, only BM samples with viability >80% were included in analysis. Peripheral blood analysis was performed using a Sysmex XT-2000 blood analyzer (Sysmex Corp.).

**Flow cytometry, sorting, and immunofluorescence.** Cell analysis was performed with a FACSCanto II flow cytometer using FACSDiva 6.1.3 software (BD) or FACS LSR II flow cytometer. BM cells were harvested from femurs by repeatedly flushing with PBS supplemented with 5% FBS (Life Technologies). Mouse spleens were homogenized, and single-cell suspensions were prepared after erythrocyte lysis with red blood cell lysis buffer (eBioscience). The analysis of BM and spleen cellularity was performed with MACSQuant (Miltenyi Biotec). The antibodies used for cell surface phenotype analysis are shown in Supplemental Table 1. Brilliant Violet 510 Streptavidin (BioLegend) was used for the detection of biotinylated antibodies. Cell viability was assessed using Hoechst 33258 (Life Technologies). Data analysis was performed using FlowJo (Tree Star) software.

**Cell sorting.** Cell sorting was performed using a FACSAria cell sorter (BD). Enrichment of Lin− cells was performed prior to cell sorting by negative magnetic selection using a MidiMACS Separator (Miltenyi Biotec). For this reason, cells were incubated with a lineage cocktail of biotinylated antibodies (Biotin-Conjugated Mouse Lineage Panel, BD Pharmingen) and subsequently with anti-Biotin MicroBeads (Miltenyi Biotec). To isolate endothelial cells, femurs and tibiae were crushed with a mortar and incubated for 30 minutes with collagenase type I (3 mg/ml, Life Technologies) prior to enrichment of Lin− cells. Endothelial cells were characterized as artheroslerotic endothelial cells (CD45 Lin− CD31 Sca1+) and sinusoidal endothelial cells (CD45 Lin CD31 Sca1−). Cells were further characterized according to Vcam1 expression, as Vcam1+ and Vcam1−.

For isolation of CXCL12-expressing niche cells, femurs were flushed and incubated with 1 U ml−1 liberase (Roche Applied Science) and 12 μm l−1 DNase I (Sigma-Aldrich) in HBSS for 30 minutes at 37°C. Different subsets of CXCL12-expressing niche cells were identified by staining digested BM suspensions from CXCL12−/− reporter mice with biotinylated antibodies against TER-119 and CD45 (clone 30-F11; eBioscience), followed by incubation with streptavidin conjugated to DyLight 405 (Jackson ImmunoResearch Laboratories Inc.) and an anti-CD31 antibody (clone 390; eBioscience). High or intermediate levels of GFP in the CD45/TER119-negative fraction were used.
to discriminate CAR cells (CD45 TER119 CD31 GFP®), CXCL12+ endothelial cells (CD45 TER119 CD31 GFP®), and CD31 GFP+ mesenchymal stromal cells (CD45 TER119 CD31 GFP®) (33, 34). Cells were sorted on a FACSARia (BD Biosciences) to >95% purity.

For immunofluorescence, tibiae were collected and fixed for 2 hours in 4% paraformaldehyde. Bone decalcification was performed for 48 hours in OsteoSoft (Merck Millipore), and bones were embedded in optical cutting temperature compound (OCT; Tissue-Tek). Endothelial cell staining in decalcified bones was performed using an APC-conjugated anti-CD31/PECAM-1 (clone 390; ebioscience) antibody and Alexa Fluor 694-conjugated anti-CD144/VE-cadherin (clone BV13; BioLegend), as described previously (7). In other experiments endothelial cell staining in bones was performed with a rat anti-mouse CD144/VE-cadherin (clone 1D4.1; BD Biosciences) antibody, followed by a goat anti-rat antibody conjugated with Alexa Fluor 647 (Thermo Fisher Scientific). In another set of experiments, staining of the vascular lumen was performed by administering an Alexa Fluor 594-conjugated Isolectin GS-IB4 from Griffonia simplicifolia (Thermo Scientific), as described (75). Del-1 staining was performed with a rabbit anti-mouse Del-1 (Proteintech, catalog 12580-1-AP), followed by a goat anti-rabbit antibody conjugated with Alexa Fluor 488 (Thermo Scientific). Nuclear staining was performed with DAPI. Images were acquired with a Zeiss LSM 510 confocal microscope, an IX83 microscope equipped with a Yokogawa CSU-X1 spinning disc confocal scanner (Olympus), or a Nikon Eclipse Ni-E automated upright fluorescence microscope (Nikon).

Cell cycle analysis. For cell cycle analysis, BM cells were stained for cell surface markers as described in the previous paragraph, fixed using fixation/permeabilization buffer (eBioscience), and stained with anti–Ki-67 antibody. After washing, cells were stained with DAPI (Molecular Probes) and analyzed by flow cytometry.

Proteins. Del-1–Fc generation has been previously described (30). An RGE point mutant of Del-1–Fc, in which Glu (E) was substituted for Asp (D) in the RGD motif of the second EGF repeat (Del-1[RGE]–Fc), was constructed by site-directed mutagenesis (QuickChange II mutagenesis kit; Agilent). Expression and purification were performed as described (30).

Adhesion assay. Adhesion was performed as previously described with modifications (29, 76, 77). Nunc-Immuno MicroWell 96-well solid plates (Sigma-Aldrich) were coated overnight at 4°C with Del-1–Fc, Del-1[RGE]–Fc, or recombinant human IgG1 Fc (Fc-control; R&D Systems) (10 µg/ml each), as described under Adhesion assay. MyeloCult medium supplemented with stem cell factor (20 ng/ml, PeproTech) was used for the incubation of LSK cells. Isolated LSK cells were incubated for 3 hours to study the mRNA levels of Ccnd1. To inhibit ILK, LSK cells were treated with the ILK inhibitor Cpd22 (2 µM; Merck Millipore), whereas DMSO was used as control.

In vitro differentiation assay. Sorted LT-HSCs (500 cells per well), MPPs (1,000 cells per well), or CMPs (1,000 cells per well) were cultured in suspension culture 96-well plates (Greiner Bio-One) in MyeloCult medium supplemented with stem cell factor (20 ng/ml, PeproTech). Del-1–Fc, Del-1[RGE]–Fc, or recombinant human IgG1 Fc (Fc-control; R&D Systems) (500 ng/ml each) were added in the cultures. To inhibit β3 integrin, LT-HSCs were preincubated for 15 minutes before the addition of Del-1–Fc or Fc-control with cilengitide (25 µg/ml; Selleck Chemicals) or RGD control peptide (25 µg/ml; Enzo Life Sciences, GRADSP). After 48 hours or on day 7, the frequency of progenitor cells was analyzed by flow cytometry. The images from the colonies were acquired using a Zeiss Observer.Z1 inverted microscope.

Real-time PCR. For RNA isolation, cells were directly sorted into RNA isolation buffer, and RNA isolation was performed using an RNasy Plus Micro Kit (QIAGEN), according to the manufacturer’s instructions. RNA from the cBM and the endosteal region was collected using TRI reagent (Molecular Research Center Inc.), as previously described (5). Briefly, after repeatedly flushing the femurs into PBS supplemented with 5% FBS, for the isolation of cBM cells, the bones were flushed twice with TRI reagent to collect RNA from the endosteal region. RNA was reverse-transcribed with the iScript cDNA Synthesis Kit (Bio-Rad). Quantitative PCR was performed by using the SsoFast EvaGreen Supermix (BioRad) and gene-specific primers in a CFX984 Real-Time PCR Detection System (Bio-Rad). Relative mRNA expression levels were calculated according to the ΔΔCt method upon normalization to 18s or β2m for mouse samples and GAPDH for human samples.

CFU assay. CFU assay was performed using MethoCult GF M3434 medium (Stemcell Technologies), according to the manufacturer’s instructions. Quantification of colony formation was performed using a STEMvision Instrument (Stemcell Technologies).

G-CSF measurement. G-CSF was measured in EDTA-plasma using a Mouse G-CSF DuoSet ELISA (R&D Systems), according to the manufacturer’s instructions.

Statistics. All data are presented as mean ± SEM. Mann-Whitney U test was used for the comparison of two groups. In vitro comparison of matched groups was performed with a paired 2-tailed Student’s t test. For comparisons of multiple groups, 1-way ANOVA followed by Holm-Sidak’s multiple comparisons test was used. All statistical analysis was performed using GraphPad Prism 6 (GraphPad Software Inc.). Significance was set at P < 0.05.

Study approval. Animal experiments were approved by the Landesdirektion Sachsen, Germany; the Animal Care and Ethics Committee of the CNIC and regional authority of Madrid; and the Institutional Animal Care and Use Committee of the University of Pennsylvania. Primary human BM-derived MSCs (70, 72-74) were derived from BM aspirates from healthy donors after written informed consent was received and upon approval by the Ethics Committee of the Technische Universität Dresden.

Author contributions. IM designed the project, performed experiments, analyzed and interpreted data, and wrote the manuscript; LSC performed experiments and analyzed data; RPS performed experiments...
and analyzed and interpreted data; IK, ME, TK, MT, AZ, KR, TM, TG, PS, and MDS performed experiments; KH and BW (Penn) generated critical reagents and interpreted data; MvB, MW, and MB provided human MSCs and interpreted data; PV, TT, and AH interpreted data; BW (Dresden) designed experiments, interpreted data, supervised research, and edited the manuscript; GH supervised research, interpreted data, and edited the manuscript; TC conceived and designed the project, interpreted data, supervised research, and wrote the manuscript.

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