GPR182 is an endothelium-specific atypical chemokine receptor that maintains hematopoietic stem cell homeostasis

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G protein-coupled receptor 182 (GPR182) has been shown to be expressed in endothelial cells; however, its ligand and physiological role has remained elusive. We found GPR182 to be expressed in microvascular and lymphatic endothelial cells of all organs and to bind with nanomolar affinity the chemokines CXCL10, CXCL12, and CXCL13. In contrast to conventional chemokine receptors, binding of chemokines to GPR182 did not induce typical downstream signaling processes, including G-protein-mediated signaling or β-arrestin recruitment. GPR182 showed relatively high constitutive activity in regard to β-arrestin recruitment and rapidly internalized in a ligand-independent manner. In constitutive GPR182-deficient mice, as well as after induced endothelium-specific loss of GPR182, we found significant increases in the plasma levels of CXCL10, CXCL12, and CXCL13. Global and induced endothelium-specific GPR182-deficient mice showed a significant decrease in hematopoietic stem cells in the bone marrow as well as increased colony-forming units of hematopoietic progenitors in the blood and the spleen. Our data show that GPR182 is a new atypical chemokine receptor for CXCL10, CXCL12, and CXCL13, which is involved in the regulation of hematopoietic stem cell homeostasis.

Significance

G protein-coupled receptors (GPCRs) are important regulators of cellular and biological functions and are primary targets of therapeu tic drugs. About 100 mammalian GPCRs are still considered orphan receptors because they lack a known endogenous ligand. We report the deorphanization of GPR182, which is expressed in endothelial cells of the microvasculature. We show that GPR182 is an atypical chemokine receptor, which binds CXCL10, 12, and 13. However, binding does not induce downstream signaling. Consistent with a scavenging function of GPR182, mice lacking GPR182 have increased plasma levels of chemokines. In line with the crucial role of CXCL12 in hematopoietic stem cell homeostasis, we found that loss of GPR182 results in increased egress of hematopoietic stem cells from the bone marrow.

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The authors declare no competing interest.

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chemokines and transports them across endothelial cells or, when expressed on erythrocytes, buffers chemokine levels in the blood (18). ACKR2 functions as a scavenger receptor by binding several C-C motif chemokine ligand (CCL) chemokines and plays various roles in the immune system (19). ACKR3 only binds C-X-C motif chemokine ligand 11 (CXCL11) and CXCL12 and controls the CXCL12–CXCR4 signaling axis by direct interaction with CXCR4 and by scavenging CXCL12 (20, 21). ACKR4 binds the conventional chemokine receptor ligands CCL19, CCL21, CCL25, and CXCL19 (18, 19).

Here, we show that GPR182 functions as an atypical chemokine receptor for CXCL10, CXCL12, and CXCL13 and that it is involved in preventing hematopoietic stem cell egress from bone marrow.

Results

To analyze the in vivo expression of GPR182, we generated a bacterial artificial chromosome (BAC)–based transgenic mouse line expressing the monomeric red fluorescent protein mCherry under the control of the mouse Gpr182 promoter (Fig. 1A). In Gpr182Cherry mice, we observed GPR182 expression in most organs. Coo-staining with different markers confirmed expression of GPR182 in vascular endothelial cells of lungs, bone marrow, lymph nodes, Peyer’s patches, liver, and spleen (Fig. 1 and SI Appendix, Fig. S1). GPR182 was not expressed in lymphatic endothelial cells from conductive arterial vessels (SI Appendix, Fig. S1 B and C). In the liver and spleen, GPR182 was expressed in basically all sinusoidal cells as well as in endothelial cells of central veins and the hepatic artery (Fig. 1 B and C). Also, sinusoidal endothelial cells and high endothelial cells of the lymph nodes expressed GPR182 (Fig. 1 D and E), as well as endothelial cells of Peyer’s patches (Fig. 1F). In the bone marrow, endomucin-positive endothelial cells of the sinusoids express GPR182, whereas endomucin-negative endothelial cells of arteries, which are identified by staining for alpha-smooth muscle actin (αSMA), do not express GPR182 (Fig. 1G). This could be confirmed by single-cell sequencing of bone marrow endothelial cells, which showed expression of Gpr182 in cells positive for the sinusoidal endothelial cell marker stabilin2 but not for arterial markers, whereas the related receptor Ackr3 was not found to be expressed in sinusoidal endothelial cells (SI Appendix, Fig. S1D). In the small and large intestine, we found GPR182 expression mainly in endothelial cells of the lamina propria, and there was no indication for expression in epithelial cells (SI Appendix, Fig. S1 E and F). We also found GPR182 to be expressed by various lymphatic endothelial cells including those of the skin, the intestine, and lymph nodes (Fig. 1D and SI Appendix, Fig. S2).

Since the closest parologue of GPR182, the atypical chemokine receptor 3 (ACKR3, also known as CXCR7), binds CXCL11 and CXCL12, we studied binding of fluorescently labeled human CXCL12 to human GPR182. Saturation binding kinetics revealed that CXCL12 indeed bound to the receptor expressed in HEK293 cells with a dissociation constant (K$_D$) of 41 nM (Fig. 2A). We then tested whether several related human chemokines are able to compete with fluorescently labeled CXCL12 for GPR182 and found that CXCL10 was able to do so (Fig. 2B). Incubation of HEK293 cells expressing GPR182, with increasing concentrations of fluorescently labeled CXCL10, revealed that CXCL10 bound to GPR182 with a slightly higher affinity (K$_D$: 19 nM) than CXCL12 (Fig. 2C). We then systematically studied the ability of 42 different chemokines to bind to GPR182. When tested for their ability to displace fluorescently labeled CXCL10 from GPR182 expressed on HEK293 cells, most chemokines tested at 120 nM had no effect (Fig. 2D). However, in addition to CXCL12, CXCL13 and (to some degree) CCL16 and CCL19, were also able to compete with CXCL10 (Fig. 2D). We then performed systematic competition binding experiments (Fig. 2E), which revealed the highest binding affinity for CXCL13 and CXCL10 with an inhibition constant (I$_C$) of 9 and 10 nM, respectively. The two CXCL12 isoforms α and β had a slightly lower binding affinity with K$_D$ values of 31 and 19 nM, respectively. The binding affinity of CCL19 was much lower (K$_D$: 260 nM), and the affinity of CCL16 was too low to be analyzed (Fig. 2E). Very similar affinities were found for murine chemokines CXCL10, CXCL12, and CXCL13 binding to mouse GPR182 (Fig. 2F and SI Appendix, Fig. S3A).

We then tested whether binding of chemokines to GPR182 resulted in G protein–mediated signaling by testing the ability of CXCL10, CXCL12, and CXCL13 to induce Ca$^{2+}$ transients in cells expressing GPR182, their conventional chemokine receptors, and known ACKRs, together with a promiscuous G protein α-subunit. As shown in Fig. 3 A–D, CXCL10, CXCL12αβ, as well as CXCL13 induced Ca$^{2+}$ transients in cells expressing their conventional receptors CXCR3, CXCR4, and CXCR5, respectively. In contrast, no effect was seen in cells expressing GPR182 when exposed to each of the four chemokines (Fig. 3 A–D). Cells expressing ACKR3 also showed no response to CXCL12 compared to control cells (Fig. 3 B and C). Since GPR182 has recently been described to be able to interact with receptor activity–modifying proteins (RAMPs) (22), we tested whether coexpression of RAMP-1, -2, or -3 with GPR182 resulted in chemokine-induced G protein–mediated responses. However, none of the RAMPs promoted downstream signaling of GPR182 (Fig. 3 E–I). Very similar results were obtained when we determined receptor-mediated activation of Gi using the NanoBit-G protein dissociation assay (23) (Fig. 3 E–H). To study potential effects of chemokines on GPR182-dependent β-arrestin recruitment, we used the Parallel Receptorome Expression and Screening via Transcriptional Output (PRESTO-Tango) system, which measures the recruitment of protease-tagged arrestin to GPCRs (24). When we expressed the GPR182-Tango construct, we noticed that GPR182 showed strong recruitment of β-arrestin in the absence of any ligand, indicating high constitutive activity (Fig. 3F). This constitutive activity was considerably higher than that of other chemokine receptors (SI Appendix, Fig. S3B). We then tested the effect of chemokines on β-arrestin recruitment. All three chemokines increased β-arrestin recruitment through their conventional receptors, CXCR3, CXCR4, and CXCR5, whereas they had hardly any effect on β-arrestin recruitment by GPR182 (Fig. 3 J–M). However, as shown before, CXCL12 strongly induced β-arrestin recruitment through its atypical receptor ACKR3 (Fig. 3 K and L). Consistent with its strong basal and almost absent ligand-dependent recruitment of β-arrestin, GPR182, together with its ligand CXCL10, strongly internalized at 37 °C (Fig. 3M). The internalization of GPR182 was, however, not affected by the ligand (Fig. 3O) but was significantly reduced in cells with suppressed expression of β-arrestin 1 and β-arrestin 2 (Fig. 3 P and SI Appendix, Fig. S3C).

Since the in vitro data indicate that GPR182 is an atypical chemokine receptor for CXCL10, -12, and -13, which binds chemokines but does not induce downstream signaling in response to chemokine binding, we tested whether plasma chemokine levels are affected by the loss of GPR182. In GPR182-deficient mice, plasma concentrations of all three chemokines were significantly increased. While CXCL10 and CXCL12 levels were two to threefold increased, CXCL13 levels were found to be increased more than 10-fold (Fig. 4 A–C), whereas levels of related chemokines, which do not bind GPR182, were not affected (SI Appendix, Fig. S3D). To validate these findings and to test whether the acute loss of endothelial GPR182 expression would result in elevated plasma levels of chemokines, we generated inducible endothelium-specific GPR182-deficient mice (Cd54CreER$^{T2}$;Gpr182$^{	ext{flx/flx}}$ [EC-Gpr182-KO]). In contrast to control mice, induction of EC-Gpr182-KO mice resulted in a rapid increase in the plasma concentration of CXCL10, CXCL12, and CXCL13 (Fig. 4 D–F). Similar to the constitutive GPR182-deficient mice, the increase in CXCL13 levels was more pronounced than increases in CXCL10 and CXCL12 plasma levels. While the increases in CXCL10 and CXCL12 plasma levels...
levels appeared to remain stable over a time period of 20 d, CXCL13 plasma levels continued to increase after induction of endothelial GPR182 deficiency (Fig. 4 D–F).

When analyzing the architecture and composition of lymph nodes, we observed no obvious differences between control and GPR182-deficient mice (SI Appendix, Fig. S4 A–C). Peripheral blood analysis showed increased levels of neutrophils and monocytes, indicating an alteration in leukocyte generation or turnover (SI Appendix, Fig. S4 D–J). Given the expression of Gpr182 in bone marrow sinusoidal cells and the known function of CXCL12 in the regulation of hematopoietic stem cell (HSC) retention to the bone marrow niche (25), we analyzed hematopoiesis in wild-type and GPR182-deficient mice by immune phenotyping using flow cytometry. The analysis revealed a significant decrease in long-term repopulating HSCs (LT-HSCs, defined as CD150+CD48-CD34-LSK) (Fig. 5 A). In contrast, progenitor populations were not affected (Fig. 5 B and C and SI Appendix, Figs. S5 A–H and S6). No alteration in the distribution of CXCL12 was observed in the bone marrow of mice lacking GPR182 (SI Appendix, Figs. S7 and S8). To test whether an increased mobilization of HSCs from the bone marrow is mirrored by an increase in hematopoietic stem/progenitors outside the bone marrow, we determined colony-forming units

Fig. 1. GPR182 is expressed in microvascular endothelial cells. (A) Schematic of part of the BAC-based mouse Gpr182-mCherry reporter transgene, which had a total length of 234 kb. UTR: untranslated region. (B–G) Representative immunofluorescence confocal images of cryosections of the indicated organs from Gpr182-mCherry BAC transgenic mice. The mCherry signal corresponds to endogenous mCherry fluorescence. Cryosections were stained with antibodies against vascular or lymphatic endothelial markers (CD31 and ETS-related gene (ERG), as well as Proxl and Lyve1, respectively). PNAd antibody was used to specifically mark high endothelial venules from lymph nodes. mCherry was often predominantly localized in the nucleus. Shown are results of representative experiments of at least three independently performed experiments. (Scale bars, 50 μm.)
(CFUs) of hematopoietic stem/progenitors in the blood and spleen (Fig. 5D and E). CFUs were significantly increased both in the peripheral blood and in the spleen (Fig. 5D and E). This was paralleled by an increase in c-Kit+ cells in the spleen (Fig. 5F). This effect was due to GPR182 expressed in endothelial cells, since tamoxifen-induced loss of endothelial GPR182 expression in inducible endothelium-specific GPR182 knockout (EC-Gpr182-KO) mice basically recapitulated the phenotype of the constitutive GPR182 knockout (Fig. 5G–L and SI Appendix, Fig. S5 I–P).

Discussion

GPR182 has been shown to be expressed in endothelial cells, but its function has remained elusive, since the physiological ligand of the receptor is not known. In this study, we show that GPR182 binds CXCL10, CXCL12, and CXCL13 but does not couple binding of these ligands to G protein or β-arrestin activation or downstream signaling. This identifies GPR182 as an atypical chemokine receptor. The closest paralogue of GPR182, ACKR3, has an overlapping ligand binding spectrum interacting with CXCL11 and CXCL12. Thus, similar to CXCL11, which can bind to two atypical chemokine receptors, ACKR1 and ACKR3 (16, 19), CXCL12 also has two atypical receptors, which underlines the complexity of the chemokine system (26, 27). Similar to the ACKRs 1 through 4, GPR182 has an alteration in the canonical DRYLAIV motif in the second intracellular loop of conventional chemokine receptors, which is believed to be required for G protein activation and signaling (28). However, insertion of the corresponding region from CXCR4 into ACKR3 did not restore G protein signaling (29, 30). This indicates that additional properties distinguish conventional from ACKRs.

Expression of GPR182 resulted in a strong basal recruitment of β-arrestin. A constitutive interaction with β-arrestin has also
Fig. 3. GPR182 does not signal in response to ligand binding. (A–D) Effect of the indicated chemokines on [Ca\textsuperscript{2+}]\textsubscript{i}, in HEK293 cells expressing the indicated receptors together with Ca\textsuperscript{2+}-sensitive bioluminescent fusion protein (G5A) (n = 3 replicates and 3 independent experiments). (E–H) Effect of the indicated chemokines on G\textsubscript{\alpha}i activity using the G\textsubscript{\alpha}i NanoBiT assay system, as described in Methods in HEK cells transfected with the indicated human receptors (n = 3 replicates and 1 experiment). (I) Ligand-independent recruitment of \beta-arrestin by GPR182 and several other conventional and ACKRs (n = 7 replicates). (J–M) Effect of the indicated chemokines on \beta-arrestin recruitment to GPR182 using the TANGO assay system as described in Methods (n = 4 replicates and 3 independent experiments). (N) Single-cell suspension HEK293T cells expressing FLAG-tagged human GPR182 and control cells (control) were incubated at 4 °C for 1 h in the presence of 100 nM of hCXCL10-AF647 and FITC-coupled anti-FLAG antibody, followed by 3 washes and 30 min of incubation at 37 °C. Thereafter cells were fixed and stained with Hoechst dye and fluorescence analyzed by confocal microscopy. (O) Cells expressing FLAG-tagged GPR182 were incubated at 4 °C with an anti-FLAG antibody, and subsequently incubated at 37 °C in the presence or absence of hCXCL10 (100 nM). After incubation for increasing time periods, cells were stained with a secondary antibody to reveal FLAG-GPR182 at the cell surface, which was then quantified by flow cytometry. (P) Cells expressing FLAG-tagged GPR182 were, in addition, transfected with control siRNA (siControl) or siRNA against \beta-arrestin-1 and \beta-arrestin-2 (siARRB1/ARRB2). Cells were incubated at 4 °C with an anti-FLAG antibody and subsequently incubated at 37 °C for 60 min. After incubation, cells were incubated with a secondary antibody to stain FLAG-GPR182 at the cell surface. Surface staining was then quantified by flow cytometry. Data are represented as percentage of surface staining relative to basal surface staining of cells kept at 4 °C (which inhibits internalization). Shown is one representative experiment of two independently performed experiments. Shown are mean values ± SEM of one experiment.
be reported for ACKR2 and ACKR3 (31–34) but not for ACKR1 (35). In contrast to ACKR2, ACKR3, and ACKR4, which show increased interaction with β-arrestin in response to ligand binding (36–38), the constitutive β-arrestin recruitment of GPR182 was not further increased by ligand application. Consistent with a role of β-arrestin in receptor internalization, ACKR2, ACKR3, and ACKR4 show constitutive internalization (34, 37, 39, 40), which, in the case of ACKR3 and ACKR4, has been reported to be further increased by ligand binding (37, 39). This correlates with our observation that GPR182 underwent constitutive receptor internalization, which was not affected by a GPR182 ligand. Thus, we conclude that the strong β-arrestin recruitment of GPR182 leads to very efficient internalization of GPR182 and its ligands. This suggests that GPR182 may function to locally remove chemokines.

The affinity of GPR182 for CXCL12 and CXCL12 is about 10-fold and 100-fold lower than the affinity of their conventional chemokine receptors, CXCR4 and CXCR3, respectively (41, 42), whereas GPR182 binds CXCL13 with slightly higher affinity than the conventional chemokine receptor CXCR5 (43). This might explain why loss of GPR182 had a more pronounced effect on plasma levels of CXCL13 than on CXCL10 and CXCL12 plasma levels. The fact that the plasma levels of all three chemokines binding GPR182 significantly increased in mice lacking GPR182 clearly shows that GPR182 functions as a scavenger of CXCL10, CXCL12, and CXCL13. Their elevated plasma levels were still one to two orders of magnitude lower than the K_D values for their conventional chemokine receptors, CXCR2, CXCR3, and CXCR5, and it remains unclear whether the increase in systemic chemokine levels per se has any effects. It appears more likely that loss of GPR182 disturbs local chemokine activities.

Gpr182 has been reported to be expressed in the epithelium of the gastrointestinal tract and to be particularly enriched in intestinal stem cells (11). This study also reported that constitutive GPR182-deficient mice show increased proliferation of intestinal epithelial cells under stress conditions, such as regeneration after injury, in the Apem1 tumor model as well as under in vivo culture conditions, which was accompanied by ERK1/2 activation. Our Gpr182 expression reporter mouse model did not show Gpr182 expression in gastrointestinal epithelial cells, a difference to the study which we currently cannot explain. However, the reported phenotype of GPR182-deficient mice would be consistent with a role of GPR182 as an atypical chemokine receptor, which has a scavenging function for CXCL12. CXCL12 has been shown to induce activation of ERK1/2 in colon carcinoma cells as well as in crypt cells of irradiated intestinal epithelium organoids and to induce proliferation of organoid crypt cells (44–46). In addition, CXCL12 promoted intestinal epithelial recovery from radiation stress (44). In the absence of GPR182, these effects are expected to be enhanced as described by Kechele et al. (11).

Our data indicate that GPR182 is involved in HSC maintenance. This may explain the recent report that mice lacking GPR182 have slightly altered blood levels of lymphocytes and neutrophils (47), an observation we could partially confirm. Overall, the consequences of the reduction of HSCs are small. Given that CXCL12 is a key factor promoting HSC maintenance and retention in the bone marrow by activating CXCR4, which is expressed on HSCs (25, 48), the ability of GPR182 to bind and inactivate CXCL12 is the most likely mechanism by which GPR182 promotes HSC maintenance. ACKR3, which also binds CXCL12, is not involved in fetal hematopoiesis (49, 50). Its role in adult hematopoiesis is unknown, but our data show no expression of ACKR3 in sinusoidal endothelial cells, suggesting that it is not involved in GPR182-related functions. The majority of HSCs is localized close to bone marrow sinusoids, where they are in close contact with endothelial cells and so-called CXCL12-abundant reticular cells, which are also known as leptin-receptor-expressing mesenchymal stromal cells, which have a perisinusoidal localization and both express CXCL12 (25, 48, 51). This is consistent with the high expression of GPR182...
Fig. 5. Alteration of the HSC population in GPR182-deficient mice. (A–L) Quantitative analysis of hematopoietic stem and progenitor cells (A–C and G–I), hematopoietic progenitor cell colony–forming units (CFUs), in the spleen and peripheral blood (D, E, J, and K), and c-Kit+ splenic cells (F and L) in GPR182-deficient mice and littermate controls (A–F), as well as in EC-Gpr182-KO mice and corresponding controls 3 mo (G–J and L) or 3 wk (J–K) after induction of endothelium-specific GPR182 deficiency with tamoxifen. Shown are the long-term repopulating HSCs (LT-HSCs; A and G), multipotent progenitors (MPP; B and H) and lineage-negative SCA-1+ c-Kit+ (LSK; C and I) determined in both femurs of the analyzed mice; n = 7 mice (Gpr182+/+ in A–C); n = 8 mice (Gpr182−/− in A–C); n = 4 mice (Gpr182−/− in D and E); n = 3 mice (Gpr182−/− in F); n = 5 mice (Gpr182−/− in D–F); n = 5 mice (control in G–I); n = 14 mice (EC-Gpr182-KO in G–I); n = 6 mice (control and EC-Gpr182-KO in J and K); n = 5 mice [control in L]; and n = 14 mice [EC-Gpr182-KO in L]). Shown are mean values ± SEM; *P ≤ 0.05; **P ≤ 0.01; ***P ≤ 0.001; n.s., non significant. Two-tailed nonparametric Mann–Whitney U test (A–E, and G–K). Unpaired two-tailed Student’s t test (F and L).
in sinusoidal bone marrow endothelial cells and its absence in endothelial cells of bone marrow arterioles. The importance of CXCL12 expressed by these cell types for HSC homeostasis is indicated by changed HSC maintenance and retention after conditional loss of CXCL12, specifically in endothelial and perivascular stromal cells (52, 53). How CXCL12 attracts or retains HSCs to their bone marrow niches is still poorly understood. The prevailing concept suggested that CXCL12 gradients in the bone marrow attract HSCs to their niches (54). This concept has recently been challenged, and it has been proposed that, rather, local CXCL12 hotspots are involved in HSC retention (55). Our experiment, however, revealed no gross alteration in the CXCL12 distribution pattern around bone marrow sinusoids in mice lacking GPR182. Regardless of the exact mechanism by which CXCL12 promotes HSC maintenance and retention, our data show that this is promoted by the high expression of GPR182 in sinusoidal endothelial cells. Since GPR182 scaves CXCL12, it is likely to reduce the free CXCL12 concentration around sinusoidal endothelial cells. This may stabilize a CXCL12 gradient toward the HSC niche and promote HSC maintenance. Loss of GPR182 may have the opposite effect, which would explain reduced retention of HSCs in the bone marrow.

Methods

Reagents. All reagents are listed in tables added to SI Appendix, Supplemental Material and Tables S1 and S2.

Cell Lines. HEK293T cells were obtained from American Type Culture Collection. HTLA cells (a HEK293 cell line stably expressing a tTA-dependent luciferase reporter and a β-arrestin2-TEV fusion gene) were a gift from Dr. Gilad Barnea (Brown University, Providence, RI). Both cell lines were maintained in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% FBS, 2 mM L-glutamine, 5 units/mL penicillin/streptomycin. All reagents are listed in tables added to SI Appendix, Supplemental Material and Tables S1 and S2.

Ligand Assay. To determine equilibrium binding of CXCL10-AF647 or CXCL12-AF647 to GPR182, 5 × 10^4 HEK293T cells seeded in 6-well plates were transfected with a eukaryotic expression plasmid carrying the GPR182 complementary DNA (cDNA) and green fluorescent protein (GFP). 48 h later, single-cell suspension was prepared by trypsinization and filtered through a 50 μm filter. The cells were resuspended into 100 μL of binding buffer incubated at room temperature for 1 h with increasing concentration of fluorescently labeled chemokine diluted in binding buffer. Thereafter, cells were washed twice with binding buffer, fixed for 10 min with paraformaldehyde (PFA) 1% and washed twice with phosphate buffered saline (PBS). Cells were resuspended into 100 μL and transferred to a 96-well plate with black wall and transparent glass bottom. Cells were then imaged using an epifluorescence Zeiss Axio Observer microscope using AxioVision software. Pictures were analyzed with Fiji software and CXCL10-AF647 or CXCL12-AF647 fluorescent signal was quantified in GFP positive cells. Competition binding assay was performed by incubating increasing concentration of nonlabeled chemokine shortly before adding the fluorescently labeled chemokine at the indicated concentration.

For quantification of binding affinity, curve fitting and K_i were determined using one site specific binding equation in GraphPad Prism 6.07 after subtracting total binding fluorescence value by the nonspecific fluorescence binding in cells transfected with control vector. In competitive experiments, curve fitting and K_i was determined using one site fit K_i equation in GraphPad Prism 6.07 by providing the concentration (40 nM) and K_i values of fluorescently labeled Alexa647 chemokine (“hot” ligand). For analysis of CXCL10 and GPR182 Internalization. Adherent HEK293T cells were transfected with the pcDNA3 eukaryotic expression vector, encoding the N-terminally FLAG-tagged hGPR182 or with control pcDNA3 vector using Lipofectamine 2000. After 1 d, single-cell suspensions were prepared by trypsinization and seeded at a density of 5 × 10^4 cells/well of 100 μL in 96-well plates. For analysis of GPR182 internalization, cells were transfected and seeded as described above. Thereafter, cells were incubated at a concentration of 10^6 per mL with anti-FLAG-M2 antibody (Sigma [1:500]) in ice-cold binding medium for 1 h at 4°C with gentle rocking to allow for chemokine and antibody binding to the cell surface. After three washes with ice-cold binding medium, cells were resuspended in ice-cold binding medium and incubated at 4°C or 37°C for 30 min with gentle rocking followed by two washes with ice-cold binding medium. Cells were then fixed in 4% PFA (1% [vol/vol]) for 10 min at room temperature, washed, and stained with Hoechst (1 μg/mL final concentration) in PBS for 15 min. Cells were then allowed to settle down in Ibidi 8-well chamber slides and were analyzed using an inverted Leica SP8 confocal microscope.

Cell Transfection and Determination of [Ca^{2+}]_i. 2 × 10^4 cells were seeded in 96-well plates with white walls and transparent bottom and transfected with plasmids containing cDNAs encoding a calcium-sensitive bioluminescent fusion protein consisting of aequorin and GFP (56), the indicated receptors and a promiscuous G protein α-subunit using Lipofectamine 2000 (Life Technologies), as described before (57). 48 h later, cells were loaded with 5 μM coelenterazine h (Promega) in HBSS containing 1.8 mM calcium and 10 mM glucose for 2 h at 37°C. Measurements were performed using a luminometric plate reader (Flexstation 3, Molecular Devices). The area under each calcium transient was calculated by using SoftMaxPro software and expressed as area under the curve.

Gαi, Activation Assay. For determination of ligand-induced Gαi signaling, we used the recently developed NanoBiot assay (23). HEK293 cells were plated onto a 96-well plate and transfected on the same day with Galphai-LgBit, Gbeta, GgammaSmbit and the receptor (30, 6, 6, and 12 ng per well, respectively) using Lipofectamine 2000 (0.2 μl/well). After 24 h, the supernatant was aspirated and 50 μl/well of 10 μM Coelenterazine h was added and then incubated at 37°C for 1 h. The plate was then transferred to the plate reader (Flexstation 3), and luciferase activity was determined in real-time before and after addition of the indicated chemokine for 5 min. Data were expressed relative to baseline (before addition of chemokine). The processed data were plotted in GraphPad Prism 6.07 and expressed as area under the curve.

β-Arrestin Assay. To determine ligand-induced interaction of receptors and β-arrestin, we used the TANGO assay (24). The Tango plasmid library, a gift from Bryan Roth, was obtained from Addgene Kit #1000000668. HTLA cells were transfected with receptor Tango in 96-well plates with white walls and transparent bottom. Thereafter, cells were incubated for 4 h in serum-free medium and were then stimulated for 16 h with the indicated ligands diluted in sterile HBSS buffer containing 1.8 mM Ca^{2+} and 10 mM glucose. The next day, the supernatant was removed and replaced by 100 μL assay buffer containing 10% Bright-Glo reagent (Promega). A 10 min incubation period at room temperature, luminescence was counted in a plate reader and expressed as relative luminescence units.

For comparison of basal activity of TANGO-CXC3, TANGO-CXCR4, TANGO-CXCR5, TANGO-ACKR3, and TANGO-GPR182, transfected cells (in quadruplicates) were fixed with 4% PFA in PBS for 10 min at room temperature. The cells were then washed three times with PBS and blocked and permeabilized with PBS containing 0.1% Triton X-100 and 5% horse serum for 30 min at room temperature. To detect total receptor expression, the cells were then incubated with a mouse anti-FLAG M2 antibody (1:1,000) diluted in PBS containing 1% bovine serum albumin (BSA) for 1 h at room temperature. After three washes with PBS, the cells were incubated with Alexa488 conjugated anti-mouse secondary antibodies (1:500) and DAPI (1:1,000) in PBS containing 1% BSA for 30 min at room temperature. Cells were then imaged using an epifluorescence Zeiss Axio Observer or with Axio Vision software. One picture per replicate was recorded by selecting a zone containing confluent cells as visualized by DAPI staining. The mean fluorescence intensity of the FL1 (per field of view) was quantified using Imagej and corrected by subtracting the background fluorescence from untransfected cells. The results of quadruplicates were averaged, and the obtained value was used to normalize the basal TANGO bioluminescence measured in seven independent wells from the same transfection.

Analysis of CXCL10 and GPR182 Internalization. Rat hepatocyte cell line (23) was transfected with receptor Tango in 96-well plates with white walls and transparent bottom. Thereafter, cells were incubated for 4 h in serum-free medium and then washed for 1 h with complete medium. Cells were resuspended in ice-cold binding medium, cells were resuspended in ice-cold binding medium and incubated 1 h at room temperature in complete medium. Cells were rinsed with binding buffer (hanks balanced salt solution [HBSS] pH 7.4, SI Appendix, Supplemental Material) and dispensed in V-bottom 96-well plate. Cells were resuspended in 100 μL binding buffer incubated at room temperature for 1 h with increasing concentration of fluorescently labeled chemokine diluted in binding buffer. Thereafter, cells were washed twice with binding buffer, fixed for 10 min with paraformaldehyde (PFA) 1% and washed twice with phosphate buffered saline (PBS). Cells were resuspended into 100 μL and transferred to a 96-well plate with black wall and transparent glass bottom. Cells were then imaged using an epifluorescence Zeiss Axio Observer microscope using AxioVision software. Pictures were analyzed with Fiji software and CXCL10-AF647 or CXCL12-AF647 fluorescent signals was quantified in GFP positive cells. Competition binding assay was performed by incubating increasing concentration of non-labeled chemokine shortly before adding the fluorescently labeled chemokine at the indicated concentration.

For quantification of binding affinity, curve fitting and K_i were determined using one site specific binding equation in GraphPad Prism 6.07 after subtracting total binding fluorescence value by the non-specific fluorescence binding in cells transfected with control vector. In competitive experiments, curve fitting and K_i was determined using one site fit K_i equation in GraphPad Prism 6.07 by providing the concentration (40 nM) and K_i values of fluorescently labeled Alexa647 chemokine (“hot” ligand).
were washed three times with ice-cold PBS before capture and analysis of 2 × 10^5 total events by flow cytometry using a FACS Canto flow cytometer (BD Biosciences). The total amount of receptor on the cell surface was determined by median fluorescence intensity and expressed as the percent of basal cell surface GPR182 expression in cells maintained at 4 °C without chemokine treatment.

For analysis of the effect of α-actinin knockdown on α-actinin internalization, adherent HEK293T cells were transfected with 20 nM control small interfering RNA (siRNA) or a mix of 10 nM siRNA targeting ARRB1 and 10 nM of siRNA targeting ARRB2 using lipofectamine RNAiMAX (Thermofisher) following manufacturer’s instructions. Two days after siRNA transfection, cells were transfected with the pCDA3 eukaryotic expression vector encoding the N-terminally FLAG-tagged human GPR182 or with control pCDA3 vector using Lipofectamine 2000. At 72 h after siRNA transfection, 10^6 cells per mL were incubated with anti-FLAG-M2 antibody (Sigma [1:500]) in ice-cold binding medium for 1 h at 4 °C with gentle rocking. After three washes with ice-cold binding medium, cells were incubated for 60 min at 37 °C with gentle rocking or were maintained at 4 °C for 30 min at 4 °C with gentle rocking. Finally, cells were washed three times with ice-cold PBS before capture and analysis of 2 × 10^5 total events by flow cytometry using a FACS Canto flow cytometer (BD Biosciences). The total amount of receptor on the cell surface was determined by median fluorescence intensity and expressed as the percent of basal cell surface GPR182 expression in cells maintained at 4 °C. Knockdown efficiency of α-actinin was determined by western blotting using standard methods.

Detection of Chemokine Levels. After puncture of the facial vein, mouse blood was collected into EDTA-coated tubes (Microvette, NC0973120), placed on ice, and spun down at 2,000 × g for 10 min at 4 °C. Plasma was then collected and frozen at −80 °C until analysis. Plasma cytokine and chemokine levels were determined by MILLIPLEX MAP magnetic bead-based multi-analyte panel mouse cytokine/chemokine 25 Plex (McytOMAX-70K-PMX) or a custom designed Mouse Magnetic LumineX Assay 3plex panel for CXCL10, CXCL11, and CXCL13 (R&D). Samples were analyzed using the MAGPIX system and MILLIPLEX Analyst 5.1 software (Merck Millipore).

Lineage and Progenitor Staining of Bone Marrow Cells. Both femurs were collected from each mouse for single mouse analysis and cleaned to remove excess muscle tissue and tendons. Bone marrow cells were extracted by flushing out from bones in 3 mL cold 1× PBS by using a 5 mL syringe applied with needle. For enrichment of bone marrow mononuclear cells (BMMNCs) the resuspended cells (3 mL) were carefully placed on top of 3 mL Histopaque 1083 (Sigma) medium containing heparin (100 mg/mL, Merck). Spleens were subsequently collected and placed in Iscove’s modified Dulbecco’s medium (IMDM) medium supplemented with 2% FBS (StemCell Technologies). Spleens were minced with surgical scissors and smashed on a 40 μm cell-strainer in order to obtain single-cell suspension. Spleenocytes were washed twice with IMDM medium containing 2% FBS, resuspended in IMDM medium, and counted using a Neubauer hemocytometer. Whole blood was treated with nine volumes of ammonium chloride solution (StemCell Technologies). Thereafter, blood cells were washed twice with IMDM medium and counted again. 2 to 3 × 10^6 spleenocytes and erythrocyte-depleted blood cells were resuspended into Methocult medium and plated in Smart-6-well plates (StemCell Technologies). The number of colonies formed was assessed after 12 d of incubation at 37 °C, 5% CO2, and ≥ 95% humidity.

Blood Analysis. For complete blood count analysis, 10 to 12 wk old mice were sacrificed by CO2, and blood was immediately collected by cardiac puncture and transferred to EDTA-coated microtubes. The collected blood as well as blood smears were analyzed by IDEXX laboratories.

Animal Models. All mice were backcrossed onto a C57BL/6J background at least 8 to 10 times, and experiments were performed with littermates as closely as possible. Male and female mice were used at 8 to 12 wk old. Mice were housed under a 12-h light–dark cycle, with free access to food and water under specific pathogen-free conditions unless stated otherwise. Mice were housed under a 12-h light–dark cycle, with free access to food and water under specific pathogen-free conditions unless stated otherwise. Transgenic mice expressing mCherry under the control of the Gpr182 promoter (Gpr182-mCherry) were generated using the BAC clone RP23-11981 from mouse chromosome 10 containing the Gpr182 gene. The coding sequence of the Gpr182gene on the BAC was replaced by a cassette carrying the mCherry cDNA, followed by a polyadenylation signal and a GFP recognition target (FRT)-flanked ampicillin resistant gene (β-lactamase) using Red/ET recombination kit (Gene Bridges). Correct targeting was verified by restriction digests and DNA sequencing. After Flp-mediated excision of the ampicillin resistant gene and linearization, the recombined BAC, which had a total length of 234 kb, was injected into pronuclei of Friend Virus NIH Jackson (FVB/N) oocytes. Transgenic offspring was genotyped by PCR insertion by genomic PCRs. In total, two independent BAC transgenic lines were produced, which showed an identical mCherry expression pattern. Mice lacking GPR182 were obtained from the Knockout Mouse Project (KOMP) Repository (knockout first allele, Gpr182tm2a(KOMP)Wtsi). To generate a conditional allele of Gpr182, in which the coding sequence of exon 2 of Gpr182 is flanked by loxP sites, a cassette flanked by FRT sites was removed by crossing mice with the FLP-deleter mouse line (S8). After FLP-mediated recombination, mice were crossed with Cdhs-CreERT2 mice (S9) to obtain animals with inducible, endothelium-specific deficiency. Maintenance of the animals was in agreement with German animal welfare legislations.

Histology and Microscopy. For analysis of mCherry reporter fluorescence, mice were sacrificed by CO2 and perfused with 20 mL PBS through the left ventricle, and 5 mL PBS containing 4% PFA in PBS was perfused in PFA. 4% PFA was added in PBS to a 4 °C for 1 h (lymph nodes, Peyer’s Patches, aorta, ear skin, and part of jejunum) or for 16 h. After fixation, organs were washed at least three times with PBS and transferred to PBS containing 30% (wt/vol) sucrose at 4 °C for 24 h. Tissues were then embedded in optimal cutting temperature (OCT) compound and stored at −80°C until further processing. OCT blocks were sectioned using cryostat, and 12 μm or 20 μm sections were mounted on SuperFrost PLUS microscope slides. For aorta, ear skin, or jejunum whole-mount immunostaining, tissues were fixed as described above and washed with PBS and further processed for immunofluorescence staining.

For immunofluorescence staining, cryosections were allowed to dry for 1 h at room temperature, washed three times with PBS, and blocked/permeabilized with antibody diluent (PBS containing 0.1% Triton X-100 and 5% horse serum) for 1 h at room temperature. Then cryosections were incubated with primary antibodies diluted in antibody diluent overnight at 4 °C (SI Appendix, Supplementary Material). After three washes with PBS, sections were incubated for 1 h at room temperature with AlexaFluor-488, -594, -647 conjugated secondary antibodies (1:500) in antibody diluent containing 4’,6-diamidino-2-phenylindole (DAPI, 1:1,000). After three washes with PBS, sections were mounted in Fluoromount and covered with glass coverslips and analyzed by confocal microscopy using Leica True Confocal Scanning (TCS) SP8 or Zeiss laser scanning microscope (LSM). To perform whole-mount aortas/middle aorta, it was clear that there was similar to cryosections. For analysis of mCherry endogenous fluorescence and respective mCherry negative samples, all procedures were performed by minimizing exposure to ambient light.

Histopathological Analysis. Hematoxylin and eosin-stained sections were examined at ×200 and ×400 magnification. To perform immunohistochemistry on cryosections, sections were stained with the rabbit monoclonal anti GPR182 (Sigma [1:50]), followed by AlexaFluor-594-conjugated donkey anti-rabbit IgG (ThermoFisher Scientific [1:300]), and anti-CD31 (BD Biosciences). Coverslips were mounted with mounting medium (Cytoseal 60, ThermoFisher Scientific) and analyzed on a FV1000 ( Olympus) confocal microscope. Nuclei were stained with 4’,6-diamidino-2-phenylindole (DAPI, 1:1,000). Images were captured using appropriate excitation and emission filters. Images were analyzed using Adobe Photoshop CS5.0 and Fiji.

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GPR182 is an endothelium-specific atypical chemokine receptor that maintains hematopoietic stem cell homeostasis.

PHARMACOLOGY

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