A novel model for ectopic, chronic, intravital multiphoton imaging of bone marrow vasculature and architecture in split femurs

Mirela Bălan\textsuperscript{1,2,3} and Friedemann Kiefer\textsuperscript{1,2,*}

\textsuperscript{1}Mammalian Cell Signaling Laboratory; Max Planck Institute for Molecular Biomedicine; Münster, Germany; \textsuperscript{2}Cluster of Excellence; Cells in Motion, CiM; Münster, Germany; \textsuperscript{3}Current address: Division of Vascular Biology; Department of Medical Biochemistry and Biophysics (MBB); Karolinska Institute; Stockholm, Sweden

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Abbreviations: BM, bone marrow; BV, blood vessels; CFU, colony forming unit; DSC, dorsal skinfold chamber; DsRed, \textit{Discosoma} sp. red fluorescent protein; EC, endothelial cells; EMCN, endomucin; GFP, green fluorescent protein; HC, haematopoietic cells; MIP, maximum intensity projection; OPN, osteopontin; P, postnatal day; PI, propidium iodide; RFP, red fluorescent protein; SHG, second harmonic generation; SOC, secondary ossification center.

Creating a model for intravital visualization of femoral bone marrow, a major site of hematopoiesis in adult mammalian organisms, poses a serious challenge, in that it needs to overcome bone opacity and the inaccessibility of marrow. Furthermore, meaningful analysis of bone marrow developmental and differentiation processes requires the repetitive observation of the same site over long periods of time, which we refer to as chronic imaging. To surmount these issues, we developed a chronic intravital imaging model that allows the observation of split femurs, ectopically transplanted into a dorsal skinfold chamber of a host mouse. Repeated, long term observations are facilitated by multiphoton microscopy, an imaging technique that combines superior imaging capacity at greater tissue depth with low phototoxicity. The transplanted, ectopic femur was stabilized by its sterile environment and rapidly connected to the host vasculature, allowing further development and observation of extended processes. After optimizing transplant age and grafting procedure, we observed the development of new woven bone and maturation of secondary ossification centers in the transplanted femurs, preceded by the sprouting of a sinusoidal-like vascular network, which was almost entirely composed of femoral endothelial cells. After two weeks, the transplant was still populated with stromal and haematopoietic cells belonging both to donor and host. Over this time frame, the transplant partially retained myeloid progenitor cells with single and multi-lineage differentiation capacity. In summary, our model allowed repeated intravital imaging of bone marrow angiogenesis and hematopoiesis. It represents a promising starting point for the development of improved chronic optical imaging models for femoral bone marrow.

Introduction

Femoral bone marrow (BM) is a major site of hematopoiesis, which crucially depends on the structural composition and histological architecture of the BM. This unique cellular environment, often referred to as the niche, enables and supports BM function and directs hematopoiesis.\textsuperscript{1} A large body of evidence suggests that disturbance of this complex 3-dimensional arrangement will impair its regulatory functions and lead to the development of pathologies.\textsuperscript{2} However, direct microscopic evidence for this assumption is scarce. The intravital microscopic interrogation of the femoral BM architecture and function is severely limited by its encasing in opaque bone.

Being central for posture and locomotion, generation of direct optical access to femoral BM by surgical thinning or removal of bone is only possible for limited periods of time, that preclude the analysis of developmental and differentiation processes.\textsuperscript{3} Furthermore, the rapid environmental changes caused by surgery, such as the alteration in oxygenation status upon exposure of the bone marrow, together with the acute post-surgical trauma, may significantly alter the behavior of BM cells.
One alternative approach is imaging of the calvarial BM, which is significantly more accessible, and the extrapolation of the resulting observations to the femoral marrow. While valuable data have been obtained by this approach, it has remained controversial how representative calvarial BM is for other BM compartments. Therefore, we aimed to create a model that for the first time allows long-term direct optical imaging of femoral BM, which we refer to as chronic BM imaging.

Our approach was based on the assumption that a dorsal skin-fold chamber (DSC), which is surgically implanted on the back skin of a mouse, could be used as a long-term transplantation and observation site for a split femur, as previously femoral bone transplants have been shown to be vascularized and to grow after transplantation into a DSC. Indeed, we report here that the transplants have been shown to be vascularized and to grow after observation site for a split femur, as previously femoral bone transplantation model enabled direct observation of the changes in vascularity and cellular composition of the BM in a femoral transplant.

Results

Transplantation of a split femur into a dorsal skin fold chamber is followed by rapid vascularisation

In this study, we aimed to develop a long-term imaging model for the vasculature and tissue architecture of femoral bone marrow. To overcome the inherent barrier of the opaque femoral bone, we decided to transplant a split femur into an ectopic, optically accessible position. For the femoral bone, we choose a splitting plane parallel to the greater trochanter and the femoral head to assure ideal optical access during later imaging (Fig. 1A).

As an optically accessible location for ectopic transplantation, we elected the dorsal skin-fold chamber (DSC), which is well established and excellently tolerated by rodents. After surgical application of a DSC, we allowed the mice to recover for 2 days and then used only those animals without any signs of inflammation or post-surgical tissue damage as transplant recipients on the third day after chamber implantation (Fig. 1D). Femurs were isolated from newborn mice, split as shown (Figs. 1B, C). To stabilize the explanted femur during the splitting process and ensure perfect orientation of the splitting plane, we fixed the bone explants in a custom parallel vice during initial experiments. Subsequently, since we noted that splitting femurs with a scalpel while being held with watchmaker forceps was highly reproducible and provided indistinguishable results from the custommade vice, we exclusively settled on this faster variant. After removal of the coverslip from the DSC, the split femur was placed in the chamber with the cut surface facing a newly applied sterile coverslip (Fig. 1E). After rescaling of the chamber, the status of the transplant was recorded and animals were observed twice daily using a stereomicroscope. In more than 70 transplants, we did not observe free erythrocytes outside the vasculature immediately after transplantation (Fig. 1E). Reproducibly, we observed the accumulation of erythrocytes in the transplant, starting between 24 and 48 hrs after transplantation and covering the whole split surface within 4 days after transplantation. We also noted the onset of active perfusion of the transplant as indicated by a constant flow of erythrocytes in selected vessels within the transplant. Hematoma formation in and around the transplant was highly prominent by 4 d following transplantation (Fig. 1F). Concomitantly, we observed sprouting of blood vessels from the transplant toward the surrounding host tissue (Fig. 1G). As the newly forming vasculature elongated and ramiﬁed the hematoma-like accumulation of erythrocytes gradually disappeared (Fig. 1H), leading to the establishment of a new functional BM vasculature by 8 days after transplantation. This angiogenic process ﬁnally resulted in a new dense vascular network that covered the transplant and its immediate surroundings by 2 weeks after transplantation (Fig. 1I, asterisk). During this period of establishment of the new vasculature, starting as early as 5 days after transplantation, we injected various contrasting agents to observe vessel morphology, however, we did not detect obvious vascular discontinuities at any time point. Fluorescent dextran e.g. only slowly extravasated and was taken up by phagocytic cells in the transplant (Supplemental movie 1).

The development of split femurs after transplantation depends on donor age

To optimize our chronic BM imaging model, we tested the developmental potential and fate of split femurs from donors at different ages. We excluded adult donors on the basis of bone size and decreased age-related haematopoietic activity, and therefore tested femurs isolated from neonates between 5 and 17 days of age.

P05 femurs (isolated from 5 d old pups) were characterized by few and short trabeculae near the growth plate and the absence of a secondary ossification center (SOC) (Fig. 2A), which starts forming around P07, when the cartilage canal is formed from the perichondrium. Split P05 femurs rapidly vascularized, however, immunofluorescence analysis revealed that the transplanted BM cavity filled with newly forming woven bone, which displaced the haematopoietic compartment (Fig. 2B).Interestingly, we noted in a P05 femur recovered 12 d after transplantation the invasion of osteopontin + (OPN +) cells into the epiphysis, indicative of the first steps of residual development of a SOC.

Because also P10 split femurs suffered from extensive ossification of the BM cavity after transplantation (Fig. S1), we analyzed the behavior of older P17 split femurs, in which the SOC is more developed, occupying most of the epiphysis, and the trabecular bone has extended into the primary ossification center (Fig. 2C). Two weeks after transplantation, we observed persistence of an open BM cavity alongside development of trabecular bone (Fig. 2D). Overall the appearance of a transplanted P17 split femur was similar enough to a P31 femur (Fig. 2E) therefore we decided to use P17 for further transplantation experiments. We
noted that the expansion of trabecular bone had progressed since the time of transplantation, however, lagged behind the age matched control (Fig. S2). BVs were present in the marrow cavity although they were less densely packed and had a larger diameter.

The newly formed vascular network originates largely from the femoral vasculature

The observation of sprout formation from the edge of the transplant suggested that the newly forming vascular network may have largely originated from the transplant. To distinguish
the contribution of transplant and host to neo-vessel formation, we took advantage of genetically labeled mouse lines. We transplanted split femurs from mice expressing a LifeAct-GFP transgene into recipients, in which the endothelial and a large part of the myeloid compartment were labeled by Tie2-Cre driven tdRFP expression from the Rosa26 locus.

In this constellation, most of the dense vascular network overlaying the transplant originated from the transplant (Fig. 3A). While it was amply surrounded by tdRFP+ cells, we noted a remarkable absence of tdRFP+ BVs either in, overlaying or in close proximity to the transplanted femur (Fig. 3B). Newly sprouting GFP+ BVs arose from the dense vascular plexus in the marrow (Fig. 3C), crossed the bone shaft (Fig. 3C, dashed white lines) and then split into smaller BVs, which ultimately created a dense network which was functionally connected with the host environment (Fig. 3C, asterisk). In few instances, we were able to observe instances of anastomosis between transplant and host-derived endothelial cells (Figs. 3, D–F).

Newly forming vessels bear morphologic resemblance to the femoral vasculature

Different vascular beds show distinct functional characteristics of the endothelium (Figs. 4A and F). The finding that the overwhelming majority of newly formed vessels arose by sprouting from the transplant raised the question if these might recapitulate aspects of the bone marrow vasculature. Presently, there are no specific surface markers known that would be unique to BM sinusoidal vessels, hence we had to base our analysis largely on morphological criteria. To address this issue, we performed overview immunostainings of the endothelial sialomucin endomucin, which is most strongly expressed on the surface of microvascular and venous endothelial cells.14,15 We noted significant differences in branching pattern, density and morphology between the vessels within or adjacent to a transplant that had been grafted 20 days earlier, and the vasculature of the host (Fig. 4A). For this analysis, we purposefully allowed a long 20 d lapse between transplantation and observation to identify morphological properties that remained stable after vascularization.

Subsequent intravital multiphoton imaging using either quantum dots (Fig. 4C) or FITC-dextran (Fig. 4D) as a contrasting agent confirmed these differences in morphology. While host vessels underneath or within the panniculus carnosus inside the chamber were smooth, thin, with rare branching points and no vascular loops (Fig. 4D), the newly generated vessels overlaying the BM were of generally thicker and more variable diameter and appeared more chaotic, with frequent branching points that made quantification unpractical (Figs. 4 C, J). A layer analysis revealed various vascular loops overlapping in different layers of the preparation (Figs. 4G, H white arrows). Furthermore, the newly formed vessels displayed frequent vascular openings / discontinuities, possibly indicative of vessel fusion (Fig. 4 G, H, I yellow arrows) and abundant filopodia (Fig. 4 G, H, I white asterisks). Also, when we compared the median filopodial length of the transplant vessels and the normal femoral BM vasculature, we detected no significant difference (Fig. S3).

Taken together, the transplant BVs were highly reminiscent of the BVs in the BM of a P17 control neonate (Figs. 4 E, F), but very different from the host vessels of the muscular environment.
The fact that the overall distribution and diameter of the BVs in the transplant varied from bona fide femoral BVs probably resulted from the extensive sprouting and remodeling processes in the newly established plexus.

The newly formed vasculature is associated with NG2$^+$ perivascular cells

Perivascular cells fulfill important roles in stabilizing blood vessels and regulating blood flow. They also engage in reciprocal interactions with the vascular endothelium, which are important for organ specific endothelial differentiation.\(^1\)

We wondered to which extent the new vascular plexus sprouting from the transplant was invested by perivascular cells and therefore used the NG2-DsRed mouse line. In this transgenic model, NG2$^+$ perivascular and mesenchymal stromal cells are labeled by NG2 promotor-driven DsRed expression. Ten days after transplantation, we readily identified NG2$^+$ cells within the BM cavity (Figs. 5A,B) and the cortical bone of the femoral shaft (Fig. 5A). NG2$^+$ perivascular cells frequently lined blood vessels and those associated with the largest vessels displayed the tendency to wrap the vessel in parallel orientation rectangular to the direction of flow (Fig. 5B).

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NG2⁺ cells also covered the more established part of the newly formed plexus, where their cell bodies were tightly associated with the vasculature (Fig. 5C). Closer to the sprouting front, NG2⁺ cells displayed ample filopodia, were more loosely associated with and covered less of the endothelium. If the reduced coverage of the BVs by NG2⁺ cells results from mural cells following the sprouting front or if the NG2⁺ cells originate elsewhere, will need additional dynamic imaging of this transgene combination in transplanted femurs (Fig. 5D).
Split femurs provide sustainable access to image the dynamics of bone marrow angiogenic responses

As part of our characterization of the split femur model, we had defined a donor age and conditions that favored efficient reperfusion and development of an open BM cavity after transplantation of a split femur. Furthermore, we had shown that the newly forming vascular plexus was mostly derived from transplanted BM vessels and retained essential morphological characteristics of BM vasculature. We therefore tested the applicability of the split femur model for intravital imaging of the dynamic angiogenic response of vessels in the BM cavity. On average DSCs remained usable for imaging over 3 weeks after transplantation before they deteriorated due to mechanical load on the skin (also see supplementary Fig. 4). In contrast to local thinning of the calvarial or femoral bone,3,4,6 the split femur allowed access at defined positions over the entire length of the BM cavity in a single imaging session, covering epiphyses, metaphyses and diaphysis. Using multiphoton microscopy, we were able to observe the transplant up to 400 μm under the level of the cover-slip, but routinely acquired stacks covering typically 200 – 340 μm in the axial dimension, independently of the imaging position. Supplemental movie 2 shows a z-stack of 271 μm depth with a 1 μm step width, acquired from a LifeAct-GFP transplant in a LifeAct-RFP host at 3.5 days after transplantation. We started stack acquisition as close as possible to the level where the cortical bone signal is present to limit observation to the marrow that is present within the confines of the transplant, and continued in depth as far as we could acquire haematopoietic cell signals. Using time lapse microscopy, we were able to visualize typical angiogenic responses like the formation of novel vessel sprouts. Supplemental movie 3 shows a single imaging session covering 6 hours of a LifeAct-GFP expressing transplant that had been implanted 4 days earlier. Typical sprouting responses with filopodia extending tip cells were visible in 2 areas of the angiogenic front (indicated by red and yellow arrows correspondingly). To orient within the transplant, we used as anatomical landmarks the bone shaft and the relative distances to the epiphyses, which are readily visualized by 2nd harmonics generation.

Myeloid progenitor cells are retained in transplanted split femurs, however, total haematopoietic activity rapidly declines after transplantation

Contrasting of the perfused vasculature by i.v. injection of fluorescent dextran impenetrable to blood vessels demonstrated that the newly formed vascular plexus was well connected to the circulation and homogeneously perfused (Figs. 4C, 5C,D). Immunostaining of cryosections of split femurs demonstrated abundant CD45+ cells within the newly forming SOC and the BM cavity, suggesting the BM vasculature was capable of retaining haematopoietic cells after transplantation (Fig. 6B). We used genetically encoded fluorescent proteins to distinguish transplant and host derived cells. Because the kinase Syk is exclusively expressed in haematopoietic cells, but not in BM endothelium17 transplanta-

Figure 5. Distribution of NG2+ cells in a transplanted split femur. (A, B) 10 days after transplantation, NG2-expressing cells marked by DsRed were prominent within the cortical bone (A) and lined BVs (demarked by white outline) without abundant ramifications of a split femur (A, B). (C) NG2+ perivascular cells (demarked by blue outline) lined BVs (demarked by white outline) distant from the sprouting edge in an apparently stable association; (D) NG2+ perivascular cells (blue outline) liaising with BVs (white outline) at the sprouting edge were characterized by multiple filopodia (asterisks) and an apparently loose association with the BVs. (A–D) MIPs of multiphoton image stacks, 4 transplants.
We noted a sharp decline in the number of haematopoietic cells during the first 3 days after transplantation to approx. 10% of the CD45+ cell present in a freshly split femur (Fig. 6D). Subsequently, the number of haematopoietic cells recovered slightly between 7 to 10 days after transplantation and then very gradually started to decline. Concomitantly, we detected a mild increase in proliferating cells (S and G2/M phase) recovered from a split femur after 7 days (Fig. 6E). This increase in cycling activity also declined until 14 days after transplantation.

We conclude from these observations that the trauma / stress of splitting and transplantation resulted in a loss of haematopoietic cells, after which the system stabilized at a new reduced level.
The fact that haematopoietic activity was retained over the entire lifespan of the transplants, prompted us to investigate, if myeloid progenitor cells were also retained within the haematopoietic compartment of split femurs. Therefore, we recovered the BM 14 days after transplantation and following preparation of single cell suspensions, we probed for the presence of myeloid progenitors by colony formation in semisolid methylcellulose cultures. We identified and enumerated colonies present after one and again 2 weeks of incubation (Fig. 6F).

Besides single lineage colonies, we were also able to recover progenitors with multi-lineage potential giving rise to erythroid cells and 3 more myeloid lineages (granulocytes, megakaryocytes and macrophages) from transplanted split femurs (Fig. 6G). The colony type distribution was well preserved in split femurs and resembled the wildtype controls more closely than the colony distribution of cells isolated from transplanted whole bones (Fig. 6F).

**Discussion**

While the overall function and general architecture of the BM are well documented, data on the dynamic behavior of different BM cell populations and their complex interrelations are scarce. Powerful new imaging techniques, including multiphoton microscopy, coupled with innovative experimental models promise to overcome the optical limitations imposed by the mineral casing of the femoral marrow. We describe here the development of a novel imaging model for femoral BM that is characterized by the fundamental criterion to allow repetitive in vivo observation of BM over broad time intervals of days to weeks, which we refer to as chronic imaging. We designed this model around the well-established dor-sal skin-fold chamber (DSC) to provide a stable, contained environment with optical access to an ectopically transplanted, split femur. We imaged this split femur transplant using multiphoton microscopy to acquire information on deeper tissue layers and to burden the sample with lower phototoxicity than that associated with conventional laser scanning microscopy. On average 200 – 300 μm tissue were imaged in the z-dimension independently of the position of imaging along the long axis of the femur.

In a DSC, ectopically transplanted split femurs re-vascularize and continue their development in an age-dependent manner

After transplantation of longitudinally split femurs into a DSC, we persistently noted a gradual erythrocyte accumulation inside the open marrow cavity, starting between 24 and 48hrs after transplantation. Erythrocytes accumulations consistently covered the whole opened marrow cavity 4 d after transplantation. Based on our observations using stereomicroscopy, intravis-tal multiphoton microscopy and immunohistology that demonstrated transplant perfusion and development of a progressively sprouting, dense vascular network uniformly covering the femur, we interpreted the presence of hematoma as a consequence of the successful connection between the ectopically transplanted femur and the host vasculature.

As the splitting process will inevitably result in damage to the BM vasculature the most obvious explanation would be that such lesions only healed after reconnection to the host circulation and initial local hemorrhages. We can however not exclude reperfusion-induced damage to the neonate femoral vasculature or the formation of erythrocyte filled lacunae due to strong hypoxia-driven, local erythropoiesis in the absence of vascular connections with the host. After 8 days the transplant vasculature was well perfused and functional as the imaging of contrasting agents did not indicate the existence of vascular lesions (Movie 1).

We hypothesized that femurs collected postnatally would connect faster and have a better chance of adapting to the new environment, compared to tissue collected from adult donors. Multiphoton and immunohistology data however revealed that neonates between P05 and P10 were not suited as donors as the marrow cavity progressively closed with new woven bone and the epiphysis was completely covered by mesenchymal cells. This closing of the BM with new bone decreased with donor age until it was no longer observed in P17 pups. Since all femurs were subjected to identical environmental factors in the DSC, the effect is most likely age-related and results from the early neonatal femur’s inability to compensate imbalances caused by ectopic transplantation leading to a random differentiation of progenitor cells toward mesenchymal and bone lineages. Equally plausible, the closing of the marrow cavity could result from progenitor cell...
differentiation aiming to regain the shielded haematopoietic microenvironment. One intermediate phase in fracture repair consists of mesenchymal cell differentiation into osteoblasts, which will fill the gap with woven bone. The age-dependent decline in fracture healing is coupled, among other factors, with a reduced number of stem and progenitor cells and a decreased self-renewal capacity of osteoprogenitors. This is consistent with the smaller surface covered by new bone when we transplanted femurs from donors of increasing ages. Presently, it is not known if the difference in bone healing decreases to this extent between postnatal days 5 and 17 in mice, but in teenagers, the geometric mean of the time required for complete femoral fracture healing increased uniformly by 0.7 wks/yr. Nevertheless for investigations aiming to study bone development and healing, younger femur transplants might be advantageous.

The split femur gives rise to a new functional vascular network that is morphologically similar to the normal bone marrow vasculature and allows intravital imaging of angiogenic responses

Genetic labeling revealed that endothelial cells (EC) contributing to newly forming BVs in the transplant were almost completely transplant derived, anastomosing with a limited number of BVs that presented host-derived markers. This observation was unexpected, since we anticipated secreted angiogenic growth factors to be released after splitting the femur and to induce a strong angiogenic host response. Though multiphoton microscopy allows deep tissue imaging, however in our model, the bone shaft obscured assessment of vascular connections made directly underneath the bone. Therefore, we can presently not comprehensively assess the extent and origin of initial vascular connections between the femur and skin.

Recently, BM EC subtypes have been described based on the expression of the markers PECAM-1 and endomucin (Emcn). The specific EC subtype likely responsible for the revascularization of the split femur is the PECAM-1 and Emcn expressing type H, localized endosteal and metaphyseal and re-establishing the BM BVs after lethal irradiation. Hypoxia was probably not the driving force behind the exuberant sprouting observed from the transplanted split femurs, as the new environment of the femoral fracture healing increased uniformly by 0.7 wks/yr. Nevertheless for investigations aiming to study bone development and healing, younger femur transplants might be advantageous.

Split femurs retain myeloid progenitor cells at a low level: a model for reactive or stressed hematopoiesis

Within the first 3 days after transplantation, we noted a rapid loss in haematopoietic cellularity contained in split femurs, followed by a stabilization or even small recovery in transplant-derived haematopoietic cell numbers that coincided with a small increase in the frequency of cycling cells. Although hematopoiesis never recovered to the initial levels, the stabilization of the haematopoietic cell complement and the fact that myeloid progenitor cells were retained in the split femur throughout the duration of the experiment suggested that a basal level of bone marrow function was preserved after transplantation.

This makes the model potentially useful to investigate regenerative or reactive stress hematopoiesis and distinct, albeit rare, homing events. Quantification of the myeloid differentiation potential in the transplants revealed a skewing toward the granulocytic lineage, compared to cells retrieved from age-matched controls. The trauma and damage of the splitting process, but also the strong angiogenic response of the transplant likely disturbed the normal BM niche, through displacement of stromal cells and changes in the ratio of EC to other stromal cells due to extensive angiogenesis. Interestingly, the differentiation profile of cells from a split femur was closer to control BM than the profile of cells retrieved from an intact transplanted femur, suggesting that oxygen availability to the haematopoietic compartment apparently was sufficient in split femurs. The presence of host cells within the transplant, revealed by genetic labeling, deserves further investigation as it could be indicative of reparatory/healing processes or even a partial repopulation of the transplant by circulating host progenitor cells.

In conclusion, we have created a novel imaging model that allows sustainable direct, which we call chronic, optical access to femoral BM (see supplemental Fig. 4S). Our approach relied on the rapid and efficient vascularization of a split femur transplanted into a dorsal skin chamber, which had been surgically implanted on the back of a mouse. In this setting, multiphoton microscopy allowed analysis of the angiogenic behavior of BM ECs. We noted a striking resemblance of vessels sprouting after transplantation and BM sinusoids, suggesting that in this model typical parameters of the BM EC are likely retained. Split femur transplants

fusion and remodeling, which transiently left transluminal endothelial pillars, before completion of the fusion/remodeling process. Sinusoids are characterized by a thin vascular wall comprised of a thin discontinuous basement membrane and incompletely covered by perivascular cells, that send projections into the BM. We used a genetic model that expressed NG2-DsRed in various cell populations including perivascular cells, and observed that their morphology differed according to their distance to the angiogenic front: the closer NG2+ cells were to the front, the more discontinuous was their coverage of the sprouting BVs and they adopted a more active phenotype with filopodia extending into the environment. At the newly-formed vascular plexus contrasting agents showed no increased vascular leakage, suggesting that the new vessels were stable, likely also due to the stabilizing effect of perivascular cells.
maintained a basal level of various haematopoietic cells of the myeloid differentiation hierarchy thus pointing toward being suitable as a model for regenerative or stress hematopoiesis. Furthermore it fosters the hope that the sustained presence of progenitor cells will ultimately allow the direct observation and analysis of cellular interactions between haematopoietic cells, either donor or host-derived and the bone marrow niche.

**Experimental procedures**

**Mice**

All animal experiments were approved by the Animal Experimentation Committee of the county of Münster and the Federal Ministry of Nature, Environment and Consumer Protection, North Rhine Westphalia. All genetic models were used on the background of: C57Bl/6 (Janvier); the strains were Syk-Cre17; LifeAct-GFP/ -RFP,31 LyEGFP,32 NG2-DsRed,33 R26R-tdRFP,34 R26-YFP,35 Tie2-Cre (Jackson Laboratory, 004128).

**Dorsal skinfold chamber implantation and femur transplantation**

The implantation procedure was performed as described in literature,12,36 with some modifications. In short, DSC implantation consisted of sandwiching the dorsal skin between 2 titanium plates and then removing a circular skin portion the size of the observation window, before closing the chamber. Femurs were split after fixation either in a custom parallel 2 titanium plates and then removing a circular skin portion (e.g., CellTrics) to obtain a single cell suspension. Cases, the resulting suspensions were passed through cell filters in PBS-T. Incubation with secondary antibodies for 4 h at 4°C confirmed by phase-contrast microscopy of May-Gr

**In vivo imaging using multiphoton microscopy**

The multiphoton system consisted of a modified upright stage Olympus, BX51WI, equipped with a Chameleon XR laser system (Coherent), optical parametric oscillator (A.P.E.-Berlin), TriMScope scan head (LaVision BioTec GmbH), GaAsP NDD detectors (Hamamatsu, H7244a-40), and an immersion XLum-PlanFl, 20 × 0.95 NA, W/IR Objective (Olympus). Images were acquired using ImSpector software (LaVisionBioTech). A Univentor 400 pump (TSE Systems) was used to control an isofuran-ane anesthesia mix of carrier gas (air enriched with oxygen, 250–300 mL/h) and anesthetic (isoflurane: induction 2–2.5%, maintenance 0.7–1.2%). Mice were immobilized in a custom-made holder that fitted the implanted DSC and allowed prevention of hypothermia by body temperature maintenance using a HAAKE DC10-K10 (Thermo) heating circulator. Eyes were covered with protection crème.

**Contrasting agents, stains and antibodies**

FITC-dextran (2 MDa, Sigma 152251), TexasRed-dextran (70 kDa life-technologies D1830). Quantum dots Macoun Red (Evident Technologies), Hoechst 33342 (Molecular Probes H3570). αCD16/32 (clone 2.4G2, BD PharMingen, 553140), αCD3e-FITC (clone 145-2C11, BD PharMingen, 553062), αLy6G-PE (clone 1A8, BD PharMingen, 551461), αF4/80 (clone CI:A31, AbD Serotec, MCA497).

**Immunofluorescence staining**

Tissue was cut into small pieces and fixed in 4% neutral buffered PFA. After washing and cryoprotection in glycerol solutions of increasing concentration (10%, 30% and 60%), samples were embedded in OCT or bone embedding medium (8% gelatine + 1% PVP). Embedded pieces were frozen in a dry ice / isopentane slurry. Thin (10 µm) and thick (50–80µm) tissue sections were rehydrated with warm PBS, and incubated in PermBlock for 2 h (1% BSA (w/v) 0.1% Tween-20 (v/v)) in PBS) for weaker block or 3% BSA (w/v) and 0.5% (v/v) Tween-20 in PBS for stronger block). Sections were incubated with primary antibodies in PermBlock overnight at 4°C, followed by 3 10 min washes in PBS-T. Incubation with secondary antibodies for 4 h at 4°C was followed by 3 washes in PBS-T, the last one containing Hoechst 33258 (1:10.000 for 30 min). After three final washes samples were mounted in Mowiol.

**Isolation of bone marrow single cell suspensions**

BM was harvested by flushing femurs with a 1 mL syringe (27 G needle) with supplemented culture medium or FACS buffer (3% BSA in PBS). Cells of transplants were gently flushed with supplemented culture medium into a 3 cm petri dish. In all cases, the resulting suspensions were passed through cell filters (e.g., CellTrics) to obtain a single cell suspension.

**Colony forming unit assay**

Murine BM cells were collected from transplanted femurs 14 days after transplantation or from age matched mice and sedimented at 300 × g (4°C). Cells were plated in 35 mm culture dishes with semi-solid culture medium comprised of IMDM containing 1% (w/v) methylcellulose, 10% heat-inactivated fetal bovine serum, 5% deionized and delipidated BSA, 1.5% IL-3 conditioned medium, 200 µg/mL iron-saturated transferrin, 25 µg/mL dipalmitolyl choline, cholesterol and oleic acid, 5 U/mL Epo, 10 µg/mL rh insulin, 12.5 ng/mL rm SCF, 10 ng/mL rh IL-6, 1 mM β-mercaptoethanol and 15% of 5637-cell conditioned medium. Type and frequency of colonies comprised of more than 50 cells were enumerated after 7 day or 14 day incubation (37°C, 5% CO2). Correct identification of colonies was confirmed by phase-contrast microscopy of May-Grünwald Giemsa stained cytosin preparations.

**Image processing**

All images were processed using either Imaris (Bitplane) or Fiji31 software. To enhance overall features, linear enhancement of brightness and contrast was performed homogeneously on whole images. 3D architecture of the new vascular
Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

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Supplemental Material
Supplemental data for this article can be accessed on the publisher’s website.

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