Lnk-dependent axis of SCF–cKit signal for osteogenesis in bone fracture healing

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The therapeutic potential of hematopoietic stem cells/endothelial progenitor cells (HSCs/EPCs) for fracture healing has been demonstrated with evidence for enhanced vasculogenesis/angiogenesis and osteogenesis at the site of fracture. The adaptor protein Lnk has recently been identified as an essential inhibitor of stem cell factor (SCF)–cKit signaling during stem cell self-renewal, and Lnk-deficient mice demonstrate enhanced hematopoietic reconstruction. In this study, we investigated whether the loss of Lnk signaling enhances the regenerative response during fracture healing. Radiological and histological examination showed accelerated fracture healing and remodeling in Lnk-deficient mice compared with wild-type mice. Molecular, physiological, and morphological approaches showed that vasculogenesis/angiogenesis and osteogenesis were promoted in Lnk-deficient mice by the mobilization and recruitment of HSCs/EPCs via activation of the SCF–cKit signaling pathway in the periosteum zone, which established a favorable environment for bone healing and remodeling. In addition, osteoblasts (OBs) from Lnk-deficient mice had a greater potential for terminal differentiation in response to SCF–cKit signaling in vitro. These findings suggest that inhibition of Lnk may have therapeutic potential by promoting an environment conducive to vasculogenesis/angiogenesis and osteogenesis and by facilitating OB terminal differentiation, leading to enhanced fracture healing.

Embryonic stem cells in the blastocyst stage have the potential to generate any terminally differentiated cells in the body; however, other adult stem cell types, including hematopoietic stem cells/progenitor cells (HSCs/HPCs), have limited potency for postnatal tissue/organ regeneration. The hematopoietic system has traditionally been considered unique among adult stem cell types, including hematopoietic stem cells/progenitor cells (HSCs/HPCs), have limited potency for postnatal tissue/organ regeneration. The hematopoietic system has traditionally been considered unique among phenotypically characterized adult stem/progenitor cells (Slack, 2000; Blau et al., 2001; Korbling and Estrov, 2003) in that it is an organized, hierarchical system with multipotent, self-renewing stem cells at the top, lineage-committed progenitor cells in the middle, and lineage-restricted precursor cells, which give rise to terminally differentiated cells, at the bottom (Weissman, 2000). Recently, Takagi et al. (2002) reported that Lnk is expressed in hematopoietic cell lineages, and BM cells of Lnk-deficient mice are competitively superior in hematopoietic population to those of WT mice. They also clarified that not only HSC/HPC numbers but also the self-renewal capacity of some HSCs/HPCs were markedly increased in Lnk-deficient mice (Ema et al., 2005). In addition, they identified the functional domains of Lnk and developed a dominant-negative Lnk
mutant that inhibits the functions of Lnk that are endogenously expressed in the HSCs/HPCs and thereby potentiates the HPCs for engraftment (Takizawa et al., 2006). Lnk shares a pleckstrin homology domain, a Src homology 2 domain, and potential tyrosine phosphorylation sites with APS and SH-2B. It belongs to a family of adaptor proteins implicated in integration and regulation of multiple signaling events (Huang et al., 1995; Takaki et al., 1997; Yokouchi et al., 1997; Li et al., 2000; Ahmed and Pillay, 2003) and has also been suggested to act as a negative regulator in the stem cell factor (SCF)–c–Kit signaling pathway (Takaki et al., 2000, 2002).

In another category of regenerative medicine, bone formation and regeneration has been extensively researched to meet clinical demand. A biologically optimal process of fracture repair results in the restoration of normal structure and function in the injured skeletal tissue. Although most fractures heal within a certain time period with callus formation that bridges the fracture gap while bone repair takes place, a large number of patients with fractures lose valuable time because of disability or confinement, leading to a loss of productivity and income. Moreover, a significant amount (5–10%) of fractures fail to heal and result in delayed union or persistent nonunion (Marsh, 1998; Rodriguez-Merchan and Forriol, 2004). Among various causes of failed bone formation and remodeling, inappropriate neoangiogenesis is considered to be a crucial factor (Harper and Kalgsbrun, 1999; Colnot and Helms, 2001). Notably, appropriate vasculogenesis by BM endothelial progenitor cells (EPCs; Asahara et al., 1997) is emerging as a prerequisite for bone development and regeneration, and there appears to be a developmental reciprocity between endothelial cells (ECs) and osteoblasts (OBs; Karsenty and Wagner, 2002). We have recently proved a pathophysiological role and contribution of murine BM-derived Sca-1+Lin- (SL) cells, HSC/EPC–enriched fraction, for bone healing (Matsumoto et al., 2008). Another group has also reported the increase of CD34+ /AC133+ cells in peripheral blood (PB) of patients with fracture, suggesting the contribution of PB EPCs to bone healing (Laing et al., 2007). However, previous studies have demonstrated that the majority of callus-formed cells in fracture were derived from the periosteum rather than from PB (Nakazawa et al., 2004), indicating a minor contribution of BM-derived cells to fracture healing. Moreover, periosteal cells, but not endosteal BM cells, have recently been shown to be competent to produce fracture callus (Colnot, 2009). Therefore, emerging the concept of enhanced osteogenesis/angiogenesis by HSCs/EPCs, one of the novel factors responsible for stem/progenitor cell mobilization from BM, that is Lnk, attracted our research interests to develop therapeutic strategy using circulating EPCs for bone fracture.

SCF has already been reported to stimulate proliferation and differentiation of HSCs (Broudy, 1997) and mobilize HSCs/EPCs into PB (Mauch et al., 1995; Takahashi et al., 1999) by binding with cKit. Thus, we have investigated the hypothesis that a lack of Lnk signaling, dependent on the SCF–c–Kit signaling pathway, enhanced the regenerative response via vasculogenesis and osteogenesis in fracture healing by HSC/EPC mobilization and recruitment to sites of fracture in Lnk-deficient mice. In our series of experiments, we showed that a negatively controlled Lnk system contributed to a favorable environment for fracture healing by enhancing vasculogenesis/angiogenesis and osteogenesis via activation of SCF–c–Kit signaling pathway, which leads to prompt recovery from fracture. In contrast, c–Kit expression was observed in several tissues and cells, including OBs (Bilbe et al., 1996).

This is the first study showing interaction between the Lnk system and fracture healing that provides a new insight into negatively controlling the Lnk system not only to promote an environment conducive to vasculogenesis/angiogenesis and osteogenesis but also to up-regulate the potential of OB terminal differentiation so that fractures can promptly heal. Therefore, negatively regulating the Lnk system has important implications for the formulation of new therapeutic strategies to enhance bone repair.

RESULTS
Pre- and post-fracture phenotypic characterization of BM and PB
We first attempted to compare the frequency of BM HSC/EPC fraction identified as c–Kit+ SL (KSL) cells at pre-fracture and 7 d after fracture in Lnk KO mice and age-matched WT mice (C57BL/6). The percentage of BM KSL fraction was significantly higher in post-fracture bone than in pre-fracture bone, regardless of mouse phenotype; however, it was significantly high in both pre- and post-fracture bone in Lnk KO mice compared with WT mice (pre-fracture: Lnk KO, 19.3 ± 0.5, WT, 5.6 ± 0.2; post-fracture: Lnk KO, 23.7 ± 0.5, WT, 8.3 ± 0.4%, respectively; P < 0.05 for Lnk KO vs. WT in both pre- and post-fracture bone; pre- vs. post-fracture bone in both Lnk KO and WT; n = 5; Fig. 1 a). Also, the number of KSL cells in 10^5 of PB mononuclear cells (MNCs) was significantly greater in post-fracture bone than in pre-fracture bone, regardless of mouse phenotype, and it was significantly great in both pre- and post-fracture PB in Lnk KO mice compared with WT mice (prefracture: Lnk KO, 1,141.4 ± 192.0; WT, 581.9 ± 97.7, post-fracture: Lnk KO, 1,467.1 ± 235.2; WT, 941.9 ± 300.0, respectively; P < 0.01 for Lnk KO vs. WT in both pre- and post-fracture; P < 0.05 for pre- vs. post-fracture in both Lnk KO and WT; n = 5; Fig. 1 b).

We next investigated which cell populations were mobilized into PB under fracture stress. Because the KSL cell population is extremely small in PB, there was no significant difference between the number and the percentage in PB at prefracture and 7 d post-fracture (unpublished data). We therefore attempted to make a comparison of the frequency of PB HSC/EPC–enriched fraction identified as cells at prefracture, 1, 4, 7, and 14 d after fracture in Lnk KO and WT mice. The percentage of post-fracture HSC/EPC–enriched fraction in PB was significantly higher, peaking at day 1 in both Lnk KO and WT mice, than that of pre-fracture (pre-fracture: Lnk KO, 19.2 ± 0.2, WT, 6.3 ± 0.5; 1 d post-fracture: Lnk KO, 51.2 ± 1.6, WT, 19.3 ± 3.1; 4 d post-fracture: Lnk KO, 53.5 ± 3.0; WT, 57.7 ± 1.1; 7 d post-fracture: Lnk KO, 43.8 ± 4.1, WT,
positive ECs were frequently observed in Tie2/LacZ Lnk KO mice, whereas the double-positive cells were rarely identified in Tie2/LacZ WT-BMT WT mice (Fig. 2 a). The number of double-positive cells was significantly increased in Tie2/LacZ Lnk KO-BMT Lnk KO mice compared with that in Tie2/LacZ WT-BMT WT mice (Lnk KO, 410.0 ± 37.6 vs. WT, 230.0 ± 56.7/mm²; P < 0.05; n = 3; Fig. 2 b). The β-gal and OC double-positive OBs were frequently observed as lining cells along with newly formed bone surface in Tie2/LacZ Lnk KO-BMT Lnk KO mice, whereas only a few double-positive cells were identified in Tie2/LacZ WT-BMT WT mice. (Fig. 2 a) The number of double-positive cells was significantly increased in Tie2/LacZ Lnk KO-BMT Lnk KO mice compared with that in Tie2/LacZ WT-BMT WT mice (Lnk KO, 195.0 ± 21.5/mm²; P < 0.05; n = 3; Fig. 2 c). These results suggest that enhanced recruitment of mobilized Tie2+ stem/progenitor cells to fracture sites contributes to vasculogenesis and osteogenesis, leading to accelerated fracture healing in Lnk KO mice.

BM-derived SL cells exhibit multilineage differentiation potential in vitro

To explore whether SL cells have multilineage differentiation potential, specifically to ECs, OBs, and adipocytes (ADs), BM-derived SL cells were cultured in differentiation induction medium for each lineage, and each marker expression was examined by real-time RT-PCR and immunocytochemistry after culture for 7 and 14 d, respectively. The mRNA expression of EC markers (CD31: 243.8 ± 7.3 vs. 372.1 ± 7.0, P < 0.0001, n = 3; Fig. 2 d), OB markers (COL1: 784.6 ± 107.8 vs. 22,667 ± 2,114, P < 0.05; n = 3; Collagen 1-A1: 784.6 ± 107.8 vs. 22,667 ± 2,114, P < 0.05; n = 3; Fig. 2 e), and AD markers (LPL: 1.4 ± 0.1 vs. 0.3 ± 0.1, P < 0.05; n = 3; PPAR-γ: 7.1 ± 0.3 vs. 6.2 ± 0.3, P < 0.0001, n = 3; Fig. 2 f) were significantly up-regulated by culture with each lineage induction medium. These lineage marker expressions were further confirmed by immunostaining for β-galactosidase (β-gal) and CD31, an EC marker, or for β-gal and osteocalcin (OC). The β-gal and CD31 double-positive ECs were frequently observed in Tie2/LacZ Lnk KO-BMT Lnk KO mice, whereas the double-positive cells were rarely identified in Tie2/LacZ WT-BMT WT mice.
Molecular evidence of enhanced angiogenesis and osteogenesis in Lnk KO mice

To explore specific gene expression differences under fracture stress between Lnk-deficient and WT mice, angiogenesis (96)– and osteogenesis (96)–related genes of the tissue at the fracture site spotted on cDNA microarrays were hybridized with biotin-labeled cDNA probes according to the manufacturer’s instructions. In angiogenesis gene array, relative expression levels of 12 genes (ANG, Fisp12, Ctgf, VEGFR[Flt1], HGF, IL-10, Gelatinase B, Mmp9, SR-A, Msr1, Restin, Rsn, BM40, Sparc, ALK-5, Tgfbr1, THBS1, and VCAM-1) of Lnk-deficient fractured mice increased by >1.5-fold, but 5 genes (Ephrin B4, Ephb4, IFNγ, NOS3, Osteopontin, Spp1, and THBS2) decreased by >2-fold compared with WT fractured mice (Fig. 3 a). In osteogenesis gene array, 24 genes (annexin A5, OC, Bglap1, Bgn, 

immunopositivities for CD31, vWF, Flk-1 (Fig. 2 g), OC (Fig. 2 h), and adiponectin (Fig. 2 i) in SL cells. The alizarin red and oil red O staining also show characteristics of OBs and ADs, respectively (Fig. 2, h and i).
expression of several key genes as a quantitative analysis. The results demonstrated a significantly higher expression of EC markers (CD31, VE-cadherin [VE-cad], and KDR/Flk1) in Lnk KO group compared with WT group (CD31: Lnk KO, 5.803 ± 0.667, WT, 1.171 ± 0.029, P < 0.01; VE-cad: Lnk KO, 7.693 ± 0.602, WT, 1.244 ± 0.006, P < 0.01; KDR: Lnk KO, 11.082 ± 1.036, WT, 1.415 ± 0.035, respectively, P < 0.01; n = 3; Fig. 3 c). The expression of bone-related

Real time RT-PCR analysis of the tissue that was RNA isolated from the peri-fracture site was also performed in the

![Figure 3. Angiogenesis- and osteogenesis-related gene expressions in sites of fracture. (a and b) A series of gene expressions detected by DNA microarray analysis in granulation tissue samples of peri-fracture sites 7 d after surgery in Lnk KO and WT mice. Red circles indicate >1.5-fold and blue circles do over 2-fold-increased blots in Lnk KO mice compared with WT mice. The selected EC-related (c) and OB-related (d) gene up-regulation in the same samples as those in DNA microarray analysis were further confirmed by quantitative real-time RT-PCR analysis. VE-cad, Flk-1 (VEGF receptor 2); Col1A1, Col1A1; and Cbf1, Cbf1. **, P < 0.01 versus WT. All data averaged with SEM from three independent experiments. All experiments were obtained from triplicated assays.

![Image of Figure 3]
enhanced neovascularization around the endochondral ossification area in Lnk KO mice compared with that in WT mice (Fig. 4 a). Neovascularization assessed by capillary density was significantly enhanced in Lnk KO mice compared with that in WT mice (Lnk KO, 1,101.8 ± 97.7 vs. WT, 628.7 ± 28.5/mm²; P < 0.01; n = 3; Fig. 4 b).

OB staining with OC (marker for mouse OB) 7 d after fracture also revealed the augmentation of osteogenesis in newly formed bone area in Lnk KO mice compared with WT mice (Fig. 4 c). Osteogenesis assessed by OB density was significantly enhanced in Lnk KO mice compared with WT mice (Lnk KO, 111.7 ± 13.2, WT, 66.7 ± 6.5/mm²; P < 0.05; n = 3; Fig. 4 d). Callus formation was monitored radiographically to evaluate fracture healing process, and the relative callus areas detected by radiography were quantified at week 1 in Lnk KO and WT group (Fig. 4 e). The relative callus area was significant larger in the Lnk KO group compared with the WT group (Lnk KO, 111.7 ± 13.2, WT, 66.7 ± 6.5/mm²; P < 0.05; n = 3; Fig. 4 d). These results indicate the enhancement of angiogenesis and osteogenesis in Lnk KO mice compared with WT mice in fracture-induced environment.

Morphological evidence of enhanced angiogenesis and osteogenesis in fractured Lnk KO mice
Enhanced angiogenesis and osteogenesis were further confirmed by immunohistochemistry. Vascular staining with CD31 (marker for mouse EC) 7 d after fracture demonstrated
fractured limb increased until 2 wk post-fracture, and then decreased slowly in KO mice. In WT mice, the blood flow showed a mild increase until 1 wk post-fracture, and then decreased. In both groups, the ratio of fractured/ intact (contralateral) blood flow significantly increased by week 1 (Fig. 5 b).

There was no significant difference in the ratio 1 h after fracture creation between the groups, whereas the ratio at weeks 1, 2, and 3 were significantly higher in Lnk KO mice compared with WT mice (0 wk: Lnk KO, 0.753 ± 0.067, WT,
Critical role of SCF–cKit signaling in enhanced SL cell mobilization from BM and recruitment to sites of fracture

Although the aforementioned results clearly demonstrated enhanced vasculogenesis/angiogenesis and osteogenesis via SL or Tie2⁺ cell mobilization into PB and recruitment to sites of fracture in Lnk KO mice, the precise mechanism remains unclear. To explore the possible mechanism, we focused on the role of the stem/progenitor cell chemokine SCF, which is a ligand for the receptor cKit, in Lnk KO mice as well as in WT mice. Accordingly, mouse (m) SCF plasma levels were measured before and after fracture at subsequent time points in both types of mice. The plasma SCF levels were decreased after fracture surgery regardless of Lnk gene deficiency in mice; however, the extent of decrease was significantly limited during the fracture healing process and returned to over the baseline level at day 14 in Lnk KO mice compared with that in WT mice (pre-fracture: Lnk KO, 263.4 ± 9.4 vs. WT, 251.0 ± 35.6 pg/ml; ns and 1 d post-fracture: Lnk KO, 194.1 ± 11.7 vs. WT, 166.8 ± 12.7 pg/ml; 4 d post-fracture: Lnk KO, 197.7 ± 7.2 vs. WT, 181.5 ± 28.9 pg/ml; 7 d post-fracture: Lnk KO, 214.7 ± 20.6 vs. WT, 190.7 ± 24.0 pg/ml; and 14 d post-fracture: Lnk KO, 284.2 ± 19.7 vs. WT, 207.3 ± 16.6 pg/ml; P < 0.01 for each comparison; n = 5; Fig. 6 a).

Next, the gain and loss of function test of SCF for SL cell mobilization in PB was performed by FACS analysis. We intraperitoneally injected 20 μg/kg mSCF, soluble SCF receptor (sKit, antagonist of SCF; Turner et al., 1995; Nakamura et al., 2004), or PBS into unfractured Lnk KO or WT mice for 5 d and examined SL cell kinetics in PB. The number of SL cells significantly increased, peaking at day 7 after SCF stimulation in Lnk KO mice, but not in WT mice, and no SL cell number increase was observed in the other treatment groups (sKit treatment in Lnk KO mice and PBS treatment in Lnk KO mice and in WT mice; prestimulation: Lnk KO-PBS, 13.0 ± 1.3; Lnk KO-SCF, 13.1 ± 0.7; Lnk KO-sKit, 15.3 ± 0.2; WT-PBS, 8.1 ± 0.9; WT-SCF, 8.7 ± 0.6 × 10⁴ cells/ml, ns, n = 5; 7 d post-stimulation: Lnk KO-SCF, 73.4 ± 4.1 vs. Lnk KO-PBS, 15.3 ± 2.5; Lnk KO-sKit, 10.0 ± 0.8; WT-PBS, 8.3 ± 0.3; and WT-SCF, 12.8 ± 2.4 × 10⁴ cells/ml, P < 0.05, n = 5; and 14 d post-fracture: Lnk KO-SCF, 39.2 ± 6.2 vs. Lnk KO-PBS, 13.6 ± 9.7; Lnk KO-sKit, 12.0 ± 0.3; WT-PBS, 7.9 ± 0.9; and WT-SCF, 10.0 ± 0.9 × 10⁴ cells/ml, P < 0.01, n = 5; Fig. 6 b). Next, to identify the synergistic effect of fracture stress and SCF stimulation on SL cell mobilization, SCF with or without sKit and PBS was injected into fractured WT mice, and sKit alone was injected into fractured Lnk KO mice in the same way. In the fractured WT mice, the number of SL cells was significantly increased, peaking at day 7 post-stimulation by SCF, but not by PBS, whereas no response of SL cell number was observed in nonfractured WT mice by SCF, and the effect of SCF on SL cell mobilization was completely cancelled by coinjection of sKit. The inhibitory effect of sKit on SL cell mobilization in PB was also observed partially in Lnk KO mice with fracture (prestimulation: Lnk KO-sKit-Fx, 16.7 ± 2.1 vs. WT-PBS-Fx, 9.0 ± 2.1; WT-SCF-Fx, 8.3 ± 0.7; and WT-SCF-sKit-Fx, 9.8 ± 0.7 × 10⁴ cells/ml, P < 0.05,

Histological evaluation with toluidine blue staining demonstrated the enhanced endochondral ossification consisting of numerous chondrocytes and newly formed trabecular bone at week 1, bridging callus formation at week 2, and complete union at week 3 in Lnk KO mice. In contrast, although a callus formation was observed at week 1, bridging callus formation was rarely found at week 2 in WT mice (Fig. 5 b). The degree of fracture healing assessed by Allen’s classification (Allen et al., 1980) was significantly higher in the Lnk KO group compared with the WT group at week 1 and 2 (1 wk: Lnk KO, 1.67 ± 0.21 vs. WT, 0.50 ± 0.22; 2 wk: Lnk KO, 2.67 ± 0.21 vs. WT, 1.50 ± 0.22, P < 0.01, n = 6; 3 wk: Lnk KO, 3.67 ± 0.21 vs. WT, 3.17 ± 0.17, ns, n = 6; Fig. 5 i). These results indicate that Lnk KO mice have a potential for prompt fracture healing evaluated by not only radiographical analysis but also biomechanical functional examination.
We further confirmed that SCF-induced stem/progenitor cell mobilization mediated enhanced angiogenesis and osteogenesis in sites of fracture by SCF–cKit signaling inhibition study in vivo. The mSCF (20 µg/kg/day), sKit (20 µg/kg/day), or PBS was injected intraperitoneally into mice with fracture for 5 d, and angiogenesis/osteogenesis was evaluated by immunofluorescent staining for CD31 and OC 7 d after surgery (Fig. 6 d). As we expected, SCF stimulation significantly increased both CD31+ capillary density (WT-SCF+/sKit−, 830.4 ± 55.7 vs. WT-SCF−/sKit−, 578.6 ± 55.2/mm², P < 0.05, n = 5; Fig. 6 e) and OC+ OB density (WT-SCF+/sKit−, 495.8 ± 52.2 vs. WT-SCF−/sKit−, 350.0 ± 25.8/mm², P < 0.05, n = 5; Fig. 6 f) around the endochondral ossification area compared with PBS injection (control: SCF−/sKit−) in WT mice, and the effect of SCF on the increase of capillary density (WT-SCF+/sKit+, 601.8 ± 60.0 vs. WT-SCF−/sKit−, 578.6 ± 55.2/mm², P < 0.05, n = 5; Fig. 6 f).

**Figure 6. Enhanced SL cell mobilization by SCF stimulation in Lnk KO mice.** (a) Serum mouse SCF levels were measured in Lnk KO and WT mice after fracture by ELISA. ††, P < 0.01; *, P < 0.05 versus Pre, respectively. (b) The number of circulating SL cells in Lnk KO and WT mice with injection of PBS (control; Lnk KO-PBS and WT-PBS), SCF (SCF, 20 µg/kg; Lnk KO-SCF and WT-SCF), or soluble SCF receptor (=c-kit ligand, sKit, 20 µg/kg; Lnk-sKit) were assessed by FACS in the indicated time course after the treatment. †† and †, P < 0.01 and P < 0.05 versus Pre, respectively. ** and *, P < 0.01 and P < 0.05 versus WT-PBS, WT-SCF, Lnk KO-PBS, and Lnk KO-sKit, respectively. (c) The number of circulating SL cells in WT mice with fracture and injection of PBS, SCF (20 µg/kg), or sKit (20 µg/kg) in Lnk KO and WT mice with fracture and sKit (WT-SCF and Lnk KO-SCF-sKit, respectively), WT mice without fracture and SCF (WT-SCF), and Lnk KO mice with fracture and sKit (Lnk KO-fx-sKit) were assessed by FACS in the indicated time course after the treatment. †† and †, P < 0.01 and P < 0.05 versus Pre, respectively. * and **, P < 0.05 and P < 0.01 versus WT-SCF-sKit, respectively. (d) Immunofluorescent staining for CD31 (green) and OC (OC, green) were performed in granulation tissue samples of peri-fracture sites in Lnk KO (left one panel) and WT (right two panels) mice 7 d after fracture. Alcian blue/orange G stained sections were also shown in parallel with immunostained sections (lower panels to each immunofluorescent staining images). Mouse phenotype with SCF (20 µg/kg) or sKit (20 µg/kg) administration (+) is indicated in the lower part of images. Dotted line indicates bone surface. Bar = 50 µm. Quantification of capillaries (e) and OBs (f). CD31 positive capillaries and OC positive OBs were counted in 3 randomly selected high power fields and averaged.**, P < 0.01 and *, P < 0.05. All data averaged with SEM from five independent experiments. All experiments were obtained from triplicated assays.
key molecule, bone morphogenetic protein (BMP)-2, was examined in Lnk KO OBs and WT OBs with SCF stimulation in the presence or absence of sKit. The significantly high BMP-2 mRNA expression was observed in Lnk KO OBs and SCF-treated WT OBs compared with WT OBs (Lnk KO-SCF−/sKit−, 2.732 ± 0.551 and WT-SCF+/sKit−, 2.576 ± 0.369 vs. WT-SCF−/sKit−, 1.189 ± 0.161, P < 0.01, n = 3 in each group) and the BMP-2 mRNA up-regulation in Lnk KO OBs and that in SCF-treated WT OBs were significantly inhibited by sKit (Lnk KO-SCF−/sKit−, 1.391 ± 0.203 vs. Lnk KO-SCF−/sKit−, 2.732 ± 0.551 and WT-SCF−/sKit−, 0.911 ± 0.136 vs. WT-SCF+/sKit−, 2.576 ± 0.369, P < 0.01, n = 3; Fig. 7 e). These results indicate that BMP-2 might be involved in the SCF–cKit signaling pathway, which is critical for OB terminal differentiation and mineralized matrix formation leading to osteogenesis enhancement in Lnk KO mice.

**SCF–cKit signaling-dependent terminal differentiation of Lnk-deficient OBs for osteogenesis**

Although there is no doubt about stimulated stem/progenitor cell recruitment to sites of fracture by SCF–cKit signaling activation in WT and Lnk KO mice, the Lnk-regulated differential potential of progenitors needed to be evaluated, especially regarding osteogenesis. We isolated mouse calvarial OBs from 3–5-d-old Lnk KO mice and WT mice, and compared the differentiation capacity in osteogenic condition medium after 7, 14, and 21 d in culture. Alkaline phosphatase (ALP) staining exhibited larger nodule formation in Lnk KO OBs than in WT OBs until day 14 (Fig. 7 a). In addition, OB mineralized matrix formation assessed by alizarin red staining in Lnk KO OBs was striking compared to that in WT OBs (Fig. 7 a). Quantitative analysis for calcium content in culture medium was also significantly high in Lnk KO OBs than that in WT OBs after 21 d in culture, (Lnk KO-SCF+/sKit−, 15.9 ± 3.5 vs. WT-SCF+/sKit−, 0.62 ± 0.1 mg/ml, P < 0.01, n = 5; Fig. 7 d). However, the number of OB CFUs (CFU-O) showed no statistical differences between the two groups at each time point (unpublished data).

Next, we investigated whether this enhancement of differentiation and mineralized matrix formation in Lnk KO OBs was also regulated by SCF signals. First, we confirmed the significantly higher mRNA expression of SCF in Lnk KO mice than that in WT OBs by real-time RT-PCR analysis (Lnk KO, 1.707 ± 0.345 vs. WT, 0.587 ± 0.111, P < 0.05, n = 3; Fig. 7 b). In WT mice, as we expected, SCF morphologically enhanced mineralized matrix formation after 21 d in culture (WT-SCF+/sKit−), and the effect of SCF on WT OBs (WT-SCF+/sKit+) and Lnk KO OBs (Lnk KO-SCF−/sKit−) was inhibited by sKit (Fig. 7 c). Calcium content in WT OBs with SCF was significantly higher than that in WT OBs without SCF (WT-SCF+/sKit+, 7.1 ± 1.4 vs. WT-SCF−/sKit−, 0.6 ± 0.1 mg/ml, P < 0.05, n = 5), and the effect was reversed by coinubcation of sKit and SCF in WT OBs (WT-SCF−/sKit−, 1.0 ± 0.3 vs. WT-SCF+/sKit+, 7.1 ± 1.4 mg/ml, P < 0.05, n = 5) and in Lnk KO OBs (Lnk KO-SCF−/sKit−, 4.8 ± 1.1 vs. Lnk KO-SCF+/sKit+, 15.9 ± 3.5 mg/ml, P < 0.01, n = 5; Fig. 7 d). We then further investigated the role of SCF–cKit signaling, focusing on an osteogenesis-related key molecule, bone morphogenetic protein (BMP)-2, was
nonadherent population of BM cells contains primitive cells able to generate both hematopoietic and osteocytic lineage cells (Olmsted-Davis et al., 2003). Zhang et al. (2003) reported that depleting a receptor of BMP in OBs caused a doubling in both OB and HSC populations in BM niche. Calvi et al. (2003) also found a parallel expansion of the HSCs when the number of OBs was increased by parathyroid hormone infusion. These findings indicate that osteogenesis and hematopoiesis/vasculogenesis closely regulated each other in terms of simultaneously resulting in accelerated bone healing in a Lnk-deficient mouse fracture model.

BM SL cells, recognized as an HSC/EPC-enriched fraction (Takahashi et al., 1999), were reported to synthesize ALP, collagen, and OC and form a mineralized matrix in culture over a decade ago (Van Vlasselaer et al., 1994). It has also been reported that BM side population cells, which contain hematopoietic repopulating cells, can also engraft in bone after transplantation (Dominici et al., 2004), and that the

Figure 7. Enhanced mineralization of OBs in Lnk-deficient mice. (a) OBs (OBs) isolated from Lnk KO and WT mice were cultured in osteogenic condition medium. Cells were assessed morphologically by ALP staining (blue) at day 14 and by alizarin red staining (red) at day 21. Upper panels show macroscopic images of whole culture dishes and lower panels show magnified images of nodule with mineralized matrix formation. (b) Real-time RT-PCR of cultured OBs for SCF mRNA expression at day 7 in Lnk KO and WT mice. Mouse heart and bone are used as positive controls for mouse endothelial and bone-related gene detection (not depicted) *, P < 0.05. Data averaged with SEM from three independent experiments. Experiments were obtained from triplicated assays. (c) The mineralized matrix formation was assessed by alizarin red staining 21 d after osteogenic culture in Lnk KO OBs with sKit (10 µg/ml) and in WT OBs in the presence of SCF (100ng/ml) with or without sKit (10 µg/ml). Upper panels show macroscopic images of whole culture dishes and lower panels show magnified images of nodule with mineralized matrix formation. Calcium content was measured by ELISA (n = 5 each; d) in culture medium and BMP2 mRNA expression (e) was analyzed by real-time RT-PCR (n = 3 each) in Lnk KO OBs with sKit (10 µg/ml) and WT OBs in the presence of SCF (100ng/ml) with or without sKit (10 µg/ml). *, P < 0.05 and **, P < 0.01. Experiments were obtained from triplicated assays.
would also be critical for fracture healing (Matsumoto et al., 2006). Indeed, cDNA microarray and quantitative real-time RT-PCR analyses also exhibited up-regulation of pro-angiogenic/-osteogenic gene expressions, which is consistent with increased blood perfusion and callus formation, in sites of fracture with recruited EPCs in Lnk-deficient mice. The series of findings suggest that both direct and indirect contribution of BM-derived EPCs to prompt and functional fracture healing with sufficient mineralization in Lnk-deficient mice.

Takaki et al. (2000, 2002) reported that Lnk acts as a negative regulator in the SCF–cKit signaling pathway. Although the role of Lnk system has gradually been clarified in the field of hematology, nothing is known about its role in skeletal biology and bone regeneration and repair. Here, we have provided evidence that enhanced HSC/EPC mobilization into PB and its recruitment to the fracture site in Lnk-deficient mice are regulated, at least in part, by SCF–cKit signaling pathway, which was proved by gain and loss of function test using SCF and sKit. However, enhanced vasculogenesis and osteogenesis in Lnk-deficient mice was shown to be significantly superior to that of SCF-treated WT mice, suggesting that another mechanism besides SCF–cKit signaling pathway is involved in the Lnk signal lacking accelerated bone healing. Indeed, molecular cDNA array analysis showed that BMPR2 mRNA expression was up-regulated more in Lnk KO mice than in WT mice. Moreover, lack of Lnk or SCF supplement in OBs induced terminal differentiation and mineralized matrix formation with increased BMP2 gene expression, which was blocked by the SCF antagonist sKit. These findings are consistent with previous studies in which cKit expression is shown in OBs and P-2 enhances OB differentiation. (Bilbe et al., 1996; Hassel et al., 2006) Collectively, it is suggested that OB differentiation and maturation is enhanced in Lnk-deficient mice via the mechanism of BMP2-involved SCF–cKit signaling pathway.
Another interesting finding in cDNA microarray data is that expression of vascular cell adhesion molecule (VCAM)-1, a differentiation-predicting marker for osteogenesis (Fukai et al., 2008), is significantly up-regulated in sites of fracture and EC-related markers in Lnk KO mice compared with WT mice. Fitau et al. (2006) reported that high expression of VCAM-1 at both the mRNA and protein level in Lnk-deficient mice was regulated in TNF-treated ECs by extracellular signal-related kinase (Erk) 1/2 pathways. In addition, Rhee et al. (2006) reported that Erk 1/2 expression was present within mesenchymal precursor cells during distraction osteogenesis and that Erk expression closely correlates with BMP 2/4 expression. Other studies also indicated that the activation of Erk 1/2 pathway in OBs related to mechanical strain and fluid flow (Jessop et al., 2002; Alford et al., 2003; Liu et al., 2008). Based on this evidence and our findings, the Erk1/2- and BMP-2/4-involved VCAM-1 signaling pathway is considered to be another mechanism for accelerated fracture healing via vasculogenesis and osteogenesis in Lnk KO mice, and further investigation will be required to clarify the entire mechanism for bone regeneration in a Lnk-deficient system.

In conclusion, our data provide novel evidence that the Lnk system acts as a negative regulator in the SCF–c–Kit signaling pathway and that Lnk deficiency modulates both vasculogenesis and osteogenesis via SL stem cell mobilization in PB and Tie2+ BM cell recruitment to sites of fracture, although Tie2+ cells are not necessarily defined as stem cells, but are likely EPCs. In addition, lack of Lnk signaling further enhances BMP2-induced OB matrix mineralization in vitro. These pathophysiological changes lead to accelerated bone healing in Lnk-deficient mice. In the field of skeletal regeneration, recently BM-derived mesenchymal stem cells or whole BMs have been used for injured bone (Petite et al., 2000) or osteogenesis imperfecta (Horwitz et al., 1999) in preclinical studies. However, there are several issues to be resolved for open reduction, such as invasiveness in cell transplantation procedure and possible complications caused by infection. The major strength of this study relies on the concept that negative control of Lnk system for bone regeneration leads to clinical feasibility by generating a Lnk-inhibitory compound. This is the first study demonstrating physiological functions of the adaptor protein Lnk in bone regeneration and suggests that inhibition of the Lnk system could be a novel therapeutic application for genetic bone diseases and bone injuries.

Isolation of Lin- BM cells. To confirm the kinetics of KSL or SL cells in BM and PB, we detected KSL or SL cells at pre-fracture and 1, 4, 7, and 14 d post-fracture by FACS analysis (n = 3 at each day).

BM cells were obtained by flushing femurs and tibias or PB cells were aspirated from the hearts of 10-wk-old Lnk KO mice or WT mice with PBS containing 5% FCS. MNCs were obtained by gradient separation onto a Ficoll Histopaque gradient. Separation of Lin- cells was performed by labeling MNCs with a lin- separation kit (BD) containing biotin-conjugated Mac1, B220, CD3e, Ter119, Ly6G, and CD45R antibodies, followed by streptavidin-conjugated magnetic beads and BD IMagnet separation before Lin- MNCs were counted.

Multilineage differentiation culture of SL BM cells. Mouse SL cells were isolated from BM MNCs by Lin- cell depletion with MACS system (Miltenyi Biotec) followed by FACS system with an anti–mouse Sca-1 antibody (FACSAria; BD), and cultured in α-modified Eagle’s medium (α-MEM; Cambrex Bio Science). After 3 wk in culture, cells (105/well) were placed in a 6-well plate and further cultured only for osteogenic and adipogenic induction.

For endothelial induction, freshly isolated SL cells were cultured in EBM-2 medium supplemented with 10% FBS and EG-M2 Bullet kit (Lonza) in Proruncept F (Sanyo Chemical, Inc.)–coated culture plate for 7 d. For osteogenic and adipogenic induction, cells were cultured in α-MEM supplemented with 10% FBS, 2 mM L-glutamine, 60 µM ascorbic acid, 10 mM β-glycerophosphate and 0.1 µM dexamethasone (Sigma-Aldrich) and α-MEM supplemented with 1 µM dexamethasone, 60 µM indomethacin, and 5 µg/ml insulin (Sigma-Aldrich) for 3 wk, respectively. After the above induction cultures, each lineage differentiation was confirmed by real-time RT-PCR fluorescence and fluorescent immunocyto staining for the indicated specific markers. Kirin, calcium deposits were detected by Alizarin red staining and formations of lipid droplets were assessed by Oil Red O staining as characteristics of OB and AD, respectively. Freshly isolated SL cells and those cultured for 3 wk in α-MEM alone were used as controls for endothelial induction and osteogenic and adipogenic induction in real-time RT-PCR analysis, respectively.

Flow cytometry studies and monoclonal antibodies. Regular flow cytometric profiles were analyzed with a FACSCalibur analyzer and CELLQuest software (BD). The instrument was aligned and calibrated daily using a four color mixture of CaliBRITE beads (BD) with FACSComp software (BD). Dead cells were excluded from the plots beads on propidium iodide (PI) staining (Sigma-Aldrich). Lineage-depleted MNCs were washed twice with HBSS containing 3.0% heat-activated FCS, and incubated with 10 µl of FcR Blocking Regent to increase the specificity of monoclonal antibodies (Miltenyi Biotec) for 20 min at 4°C, and incubated with the monoclonal antibodies for 30 min at 4°C. The stained cells were washed three times with PBS containing 3.0% FCS, and resuspended in 0.5 ml of HBSS/3%FCS/PI, and analyzed by FACScan caliber flow cytometer (BD). The following monoclonal antibodies were used to characterize the lineage-depleted MNCs: APC-conjugated anti–c–Kit (BD), FITC-conjugated anti–Sca1 (BD), IgG1-FITC isotype controls (BD), and PI (Sigma-Aldrich).

Induction of femoral fracture. All surgical procedures were performed under anesthesia and normal sterile conditions. Anesthesia was performed with ketamine hydrochloride (60 mg/kg) and xylazine hydrochloride (10 mg/kg) administered intraperitoneally. We followed Manigrasso’s model of closed femur fracture (Manigrasso and O’Connor, 2004). A lateral parapatellar knee incision on the right limb was made to expose the distal femoral condyle. A 2-mm wedge was made using a 27-gauge needle on the intercondyle of the femur and then a 0.5-mm-diam, stainless wire was inserted in a retrograde fashion. The wire was advanced until its proximal end was positioned stable to the greater trochanter and the distal end was cut close to the articular surface of the knee. A transverse femoral shaft fracture was then created in the right femur of each mouse by three-point bending. The wound was then irrigated with 10 cc of sterile saline and skin was closed in layers with 5–0

MATERIALS AND METHODS

Mice. Lnk-/- mice, whose generation and genomic cloning were described previously (Kaki et al., 2000) were backcrossed with C57BL/6 (B6-Ly5.2) >10 times and paired with age-matched WT mice as controls. Mice congenic for the Ly5 locus (B6-Ly5.1) were bred and maintained at the animal facility of Institute of Physical and Chemical Research Center for Developmental Biology, Kobe, Japan. The mice were fed a standard maintenance diet and provided water ad libitum. Male 10–12-wk-old Lnk-/- and WT mice were used in this study. The institutional animal care and use committee of the Institute of Physical and Chemical Research Center for Developmental Biology approved all animal procedures, including human cell transplantation.
nylon sutures. Postoperative pain was managed by administration of subcutaneous injection of buprenorphine hydrochloride after surgery. Unprotected weight bearing was allowed immediately after the operation. The left nonfractured femur served as a control.

20 animals were assigned to each group for radiological assessment, and every additional three to five animals in each group were assigned for each study. If the fracture produced was not a stable transverse fracture or if evidence of deep infection developed then animals were excluded from the study and replaced with another animal. Thus, six mice with comminuted fractures were replaced during the experiment. No mice developed infection, as confirmed by radiograph.

**Mouse BMT model.** Male Lnk KO mice and WT mice (C57BL6/J; CLEA Japan, Inc.) aged 6 wk were used as recipients for BMT. Transgenic mice of B6/N-TgfN (Tie-2-LacZ) [Jackom ImmunoResearch Laboratories, which were generated by backcrossing FVB/N-TgfN (Tie-2-LacZ) to C57BL6/J mice, were used as donors for the BMT. The procedure of BMT was performed as described previously, with some modifications (Li et al., 2005). In brief, the recipient mice were lethally irradiated for BM ablation with 12.0 Gy and received 5 million donor BM MNCs. At 4–6 wk after BMT, by which time the BM of the recipient mice was reconstituted, surgery for fracture induction was performed. The granulation tissue of fractured BMT mice were harvested 7 and 28 d after surgery for histological, radiographical, and biomechanical function analyses.

**Gene expression analysis via cDNA microarray.** Total RNA was obtained from tissues at the peri-fracture site at day 7 using Tri-zol (Life Technologies) according to the manufacturer’s instructions. cDNAs were synthesized using 1 µg total RNA in the presence of Superscript II and Oligo (dT)12-18 (both from Invitrogen). PCR was performed in a 20-µl reaction solution containing 2 µl 10 X PCR buffer, 150 nmol MgCl2, 10 nmol dNTP, 20 pmol primer, 1 µl 10× diluted cDNA, and 1 U RedTag DNA polymerase (Sigma-Aldrich). PCRs were run as follows: 35 cycles of 94°C for 30 s, 55°C for 30 s and 72°C for 30 s, and a final extension for 10 min at 72°C. Primer sequences are shown in the Primer size and sequence section. Nonradioactive GEArray Q series cDNA expression array filters (Mouse Angiogenesis Gene Array [MM-009] and Mouse Osteogenesis Gene Array [MM-026]; Santa Cruz Biotechnology, Inc.) were used: anti–mouse rat Flk-1 (1:100; Chemicon), anti–mouse rat CD31 (1:50; BD), anti–mouse goat vWF (1:100; Biolegend), anti–mouse rat OC (1:1,000; Cortex) to detect Tie2

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**Real-time quantitative RT-PCR analysis of RNA isolated from peri-fracture site and cultured OBs.** Total RNA was obtained from tissues of peri-fracture site at day 7 or from cultured OBs using TRIzol (Life Technologies) according to the manufacturer’s instructions. After total RNA isolation, we made single-strand cDNA using a reverse transcription kit (Invitrogen) and used it as template for real-time PCR with SYBR Green PCR Master Mix (Applied Biosystems) and gene-specific primers in an Oligo software (Takara Bio Inc.). The mean cycle threshold values from quadruplicate measurements were used to calculate the gene expression, with normalization to GAPDH as an internal control.

**Primer size and sequences.** mCD31 (224 bp): sense 5′-TCCCACCC- GAAAGCCATGAA-3′; antisense 5′-CCACGAGAAGTACTCTGCTT- TACT-3′, mVE-cad (369 bp): sense 5′-GAGCTAAGAGCCACCTCTGCTG- TA-3′, antisense 5′-TGGGCTCTTCTTTGGTCTGATG-3′, mFlk-1 (346 bp): sense 5′-TGGGCTTCCCTACTCTAATGGA-3′; antisense 5′-GAAGGCCAACAAAGCTAATGCTG-3′, mVE-cad (20 bp): sense 5′-ACGCATTCTCAGATCAA-3′; antisense 5′-AAGCATCTCCC- ACAGATTC-3′, mOC (187 bp): sense 5′-CTGACCTCAGATCCTCAA- CAGC-3′; antisense 5′-TGCTGTTGTGCCTGATCA-3′, mCOVI (107 bp): sense 5′-CAATTGTTGAGACGTTGAAAMC-3′; antisense 5′-GGTT- GGACAGTCCGTCT-3′, mChf (151 bp): sense 5′-AAGCAG- CTGAGATTTGTTGAGC-3′; antisense 5′-CCTGCGTGGTGTTAATTTCTT- GGTT-3′, mPRL (65 bp): sense 5′-AGCTGGTTGAGCAAGAATCTGTT-3′; antisense 5′-CATGTTGGTGTTGTGTTCACA-3′, mPPARy (50 bp): sense 5′-CTGCCCTCTCATGATGAA-3′; antisense 5′-AATCCCTT- GGCCCTCTGAGAT-3′, mSCF (72 bp): sense 5′-CACAAGAACAGC- AAATACAGA-3′; antisense 5′-AGACTCGGGCCCTACAAGTGA-3′, mBMP-2 (73 bp): sense 5′-TCATTATTGACCCGATACTCTTCC-3′; antisense 5′-TTGTTTATCTCATGAGGCTAACTG-3′, and mGAPDH (484 bp): sense 5′-GTGAGCGCGGTGTCAGATGATG-3′; antisense 5′-AGGCGGCACTGCAGATCC-3′.

**Tissue harvesting.** Mice were euthanized with an overdose of ketamine and xylazine. Bilateral femurs were harvested and embedded in OCT compound, snap frozen in liquid nitrogen, and stored at −80°C for histochemical staining and immunohistochemistry. Rat femurs in OCT blocks were sectioned, and 6-µm serial sections were collected on slides, followed by fixation with 4.0% paraformaldehyde at 4°C for 5 min, and then stained immediately.

**Morphometric evaluation of capillary and OB density.** Immuno- histochemical staining with anti–mouse rat CD31 (Biogenesis) for mouse EC marker or anti–mouse goat OC (Santa Cruz Biotechnology, Inc.) for mouse OB was performed. The secondary antibodies for each immunostaining are as follows: FITC-conjugated anti–rat IgG (H+L; Jackson ImmunoResearch Laboratories) for CD31 and FITC-conjugated anti–goat IgG (H+L; Jackson ImmunoResearch Laboratories) for OC. Capillary or OB density was morphometrically evaluated by histological examination of 5 randomly selected fields of tissue sections removed from segments of soft tissue in the peri-fracture site. Capillaries were recognized as tubular structures positive for CD31. OB-like cells were recognized as lining or floating cells positive for OC on new bone surface. All morphometric studies were performed by two examiners who were blinded to treatment.

**LDPI assessment.** LDPI (Moor Instrument; Wardell et al., 1993; Linden et al., 1995) was used to record serial blood flow measurements over the 3 wk post-fracture course. In these digital color-coded images, red hue indicated with maximum perfusion; medium perfusion values are shown in yellow; lowest perfusion values are represented as blue. This was done under anesthesia with the animal supine and both limbs fully fixed.

**Fluorescent immunostaining.** To detect Tie2 cell-derived neovascularization in Lnk KO mice with Tie2/LacZ, Lnk KO BM at the fracture site, double immunohistochemistry was performed with anti–mouse rat CD31 (1:50; BD) or anti–mouse goat OC (1:250; Biogenesis) and anti–rabbit β-gal (1:1,000; Cortect) to detect Tie2 cell-derived ECs or OBs. For characterization of multicell type cells in SL cells, the following primary antibodies were used: anti–mouse rat CD31 (1:50; BD), anti–mouse goat vWF (1:100; Santa Cruz Biotechnology, Inc.), anti–mouse rat Flk-1 (1:100; Chemicon), anti–mouse goat OC (1:250; Biogenesis), and anti–mouse mouse adiponectin.
Calcium content was measured in bone from link KO, WT, and fractured WT mice, and mobilization of SL cells to PB occurred intraperitoneally with 20 µg/kg SCF or PBS for 5 d in un-fractured mice. Effect of SCF stimulation on mobilization of SL cells. Concentrations of cytokines used for culture were as follows: FITC-conjugated anti-rat IgG (H+L; Jackson ImmunoResearch Laboratories) for CD31 and Fli-1, FITC-conjugated anti-goat IgG (H+L; Jackson ImmunoResearch Laboratories) for OC and vWF, Alexa Fluor 594-conjugated rabbit anti-mouse IgG (H+L) for adiponectin, and Cy3-conjugated goat anti-rabbit IgG (H+L; Jackson ImmunoResearch Laboratories) for β-gal staining. DAPI solution was applied for 5 min for nuclear staining.

Biomechanical analysis of fracture union. Biomechanical evaluation was performed with mice at week 4 after fracture in Kureha Special Laboratory. In brief, fractured femurs and the contralateral nonfractured femurs were prepared and intramedullary fixation pins were removed before the testing. The authors state that there is no conflict of interest.

Histological assessment. Histological evaluation was performed with toluidine blue staining to address the process of endochondral ossification on weeks 1, 2, and 3. The degree of fracture healing was evaluated at week 1, 2, and 3 in each group using a five point scale proposed by Allen et al. (1980). According to this classification system, grade zero indicates the formation of a pseudoarthrosis (most severe form of nonunion), grade one represents an incomplete cartilaginous union (retention of fibrous elements in the cartilaginous plate), and grade four represents complete bony union, grade three represents an incomplete cartilage uniting the fragments. Grade two represents a complete cartilaginous union (well-formed plate of hyaline cartilage) (Allen, H.L., A. Wase, and W.T. Bear. 1980. Indomethacin and aspirin: effect of indomethacin on fracture healing in young rats. Bone 5:105–111). According to this classification system, grade three represents a complete cartilaginous union (well-formed plate of hyaline cartilage) (Allen, H.L., A. Wase, and W.T. Bear. 1980. Indomethacin and aspirin: effect of indomethacin on fracture healing in young rats. Bone 5:105–111). Histological assessment was performed with toluidine blue staining to address the process of endochondral ossification on weeks 1, 2, and 3. The degree of fracture healing was evaluated at week 1, 2, and 3 in each group using a five point scale proposed by Allen et al. (1980). According to this classification system, grade four represents complete bony union, grade three represents an incomplete cartilaginous union (well-formed plate of hyaline cartilage uniting the fragments), grade two represents an incomplete cartilaginous union (retention of fibrous elements in the cartilaginous plate), and grade one represents a complete cartilaginous union (well-formed plate of hyaline cartilage). Radiographs of each animal were examined by three observers who were blinded to treatment. To evaluate the fracture healing process, callus formation was monitored radiographically and relative callus areas detected by radiography were quantified with National Institutes of Health image analysis software. In brief, fractured femurs and the contralateral nonfractured femurs were prepared and intramedullary fixation pins were removed before the imaging analysis. In brief, fractured femurs and the contralateral nonfractured femurs were prepared and intramedullary fixation pins were removed before the imaging analysis. Biomechanical analysis of fracture union. Biomechanical evaluation was performed with mice at week 4 after fracture in Kureha Special Laboratory. In brief, fractured femurs and the contralateral nonfractured femurs were prepared and intramedullary fixation pins were removed before the testing. The authors state that there is no conflict of interest.

Histochemical analysis of cell cultures. Histochemical staining for ALP was performed at day 7 and 14 using a commercially available kit (Muto-Kagaku) according to the manufacturer’s instruction. Mineralization was assessed by using alizarin red staining method. In brief, the cultures were rinsed twice with PBS, fixed in 100% ethanol for 30 min, and stained with 1% alizarin red S (Hartman Leddon Co.) in 0.28% ammonia water for 10 min at room temperature. The stained cell layers were washed, rinsed twice with distilled water, and air-dried. Matrix/intracellular calcium accumulation. Calcium content was measured by the orthocresolphthalein complexone (OCPC) colorimetric method (Sigma-Aldrich). In brief, bone density and number of trabecula in callus area were calculated with CT intensity in scanned images by single energy x-ray absorptiometry method and averaged.

ELISA assessment of plasma SFC levels. SFC plasma levels of mouse were measured at prefracture and 1, 4, 7, and 14 d post-fracture in both groups via ELISA kit (R&D Systems) according to the manufacturer’s instruction. Rat and mouse SFC and sKit were purchased from R&D Systems and supplemented concentrations of cytokines used for culture were as follows: SFC 100 ng/ml and sKit 10 µg/ml (Nakamura, 2004)

Mouse calvarial OB culture. Calvarial cells were isolated from 3-5 d-old mice using a modification of the method described by Wong and Cohn (1975). In brief, after removal of sutures, calvariae were subjected to four sequential 15-min digestions in an enzyme mixture containing 0.05% trypsin (Invitrogen) and 0.1% collagenase P (Boehringer Mannheim) at 37°C on a rocking platform. Cell fractions 2–4 were collected, and enzyme activity was stopped by the addition of an equal volume of DMEM containing 10% FCS, 100 U/ml of penicillin, and 100 µg/ml of streptomycin (Invitrogen). The fractions were pooled, centrifuged, resuspended in DMEM containing 10% FCS, and filtered through a 70-µm cell strainer. Cells were plated at a density of 105 cells/well in 35-mm culture plates in DMEM containing 10% FCS. The medium was changed 24 h later, and 3 d later cultures were fed again. At 1 wk of culture, the medium was changed to a differentiation medium (α-MEM containing 10% FCS, 50 µg/ml of ascorbic acid, and 4 mM of β-glycerophosphate) and thereafter the medium was changed every 2 d. Culture O was scored at day 7, 14, and 21 of incubation by in situ observation of plates on an inverted microscope. Effects of SCF and sKit on culture formation and mineralization were also investigated in the above assay. Purified recombinant human SFC and sKit were purchased from R&D Systems and supplemented concentrations of cytokines used for culture were as follows: SFC 100 ng/ml and sKit 10 µg/ml (Nakamura, 2004).

Statistical analysis. All values were expressed as mean ± SEM. Paired Student’s t-tests were performed for comparison of data before and after fracture. The comparisons among groups were made using the one-way analysis of variance. Post hoc analysis was performed by Fisher’s LSD test. A p-value < 0.05 was considered to denote statistical significance. It must be noted that the authors state that there is no conflict of interest.
Link-dependent axis of SCF–cKit signal for osteogenesis | Matsumoto et al.

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