Osteoporotic bones have reduced spongy bone mass, altered bone architecture, and increased marrow fat. Bone marrow stem cells from osteoporotic patients are more likely to differentiate into adipocytes than control cells, suggesting that adipocyte differentiation may play a role in osteoporosis. VEGF is highly expressed in osteoblastic precursor cells and is known to stimulate bone formation. Here we tested the hypothesis that VEGF is also an important regulator of cell fate, determining whether differentiation gives rise to osteoblasts or adipocytes. Mice with conditional VEGF deficiency in osteoblastic precursor cells exhibited an osteoporosis-like phenotype characterized by reduced bone mass and increased bone marrow fat. In addition, reduced VEGF expression in mesenchymal stem cells resulted in reduced osteoblast and increased adipocyte differentiation. Osteoblast differentiation was reduced when VEGF receptor 1 or 2 was knocked down but was unaffected by treatment with recombinant VEGF or neutralizing antibodies against VEGF. Our results suggested that VEGF controls differentiation in mesenchymal stem cells by regulating the transcription factors RUNX2 and PPARγ2 as well as through a reciprocal interaction with nuclear envelope proteins lamin A/C. Importantly, our data support a model whereby VEGF regulates differentiation through an intracrine mechanism that is distinct from the role of secreted VEGF and its receptors.
Intracellular VEGF regulates the balance between osteoblast and adipocyte differentiation

Yanqiu Liu,1 Agnes D. Berendsen,1 Shidong Jia,2 Sutada Lotinun,3 Roland Baron,3 Napoleone Ferrara,4 and Bjorn R. Olsen1

1Department of Developmental Biology, Harvard School of Dental Medicine, Boston, Massachusetts, USA. 2Department of Cancer Biology, Dana-Farber Cancer Institute, Boston, Massachusetts, USA. 3Department of Oral Medicine, Infection and Immunity, Harvard School of Dental Medicine, Boston, Massachusetts, USA. 4Genentech, South San Francisco, California, USA.

Osteoporotic bones have reduced spongy bone mass, altered bone architecture, and increased marrow fat. Bone marrow stem cells from osteoporotic patients are more likely to differentiate into adipocytes than control cells, suggesting that adipocyte differentiation may play a role in osteoporosis. VEGF is highly expressed in osteoblastic precursor cells and is known to stimulate bone formation. Here we tested the hypothesis that VEGF is also an important regulator of cell fate, determining whether differentiation gives rise to osteoblasts or adipocytes. Mice with conditional VEGF deficiency in osteoblastic precursor cells exhibited an osteoporosis-like phenotype characterized by reduced bone mass and increased bone marrow fat. In addition, reduced VEGF expression in mesenchymal stem cells resulted in reduced osteoblast and increased adipocyte differentiation. Osteoblast differentiation was reduced when VEGF receptor 1 or 2 was knocked down but was unaffected by treatment with recombinant VEGF or neutralizing antibodies against VEGF. Our results suggested that VEGF controls differentiation in mesenchymal stem cells by regulating the transcription factors RUNX2 and PPARγ as well as through a reciprocal interaction with nuclear envelope proteins lamin A/C. Importantly, our data support a model whereby VEGF regulates differentiation through an intracrine mechanism that is distinct from the role of secreted VEGF and its receptors.

Introduction

Osteoporosis, skeletal fragility associated with reduced quantity and quality of bone, is a major problem in the aging population. During aging, the balance between bone formation and resorption shifts in favor of resorption, resulting in reduced bone mass and altered bone architecture. In vitro, bone marrow mesenchymal stem cells from patients with osteoporosis are more likely to differentiate into adipocytes than osteoblasts compared with cells isolated from patients with normal bone mass (1), and osteoporotic bones have increased accumulation of adipocytes in the trabecular bone marrow space (2–5). These studies and data from mouse models of osteoporosis have prompted questions about whether increased bone marrow fat is a consequence or the cause of decreased bone mass in osteoporosis (6).

Adipocytes and osteoblasts differentiate from a common mesenchymal precursor cell. Experimental evidence indicates that a substantial degree of plasticity exists between osteoblast and adipocyte lineages (7–10). Even fully differentiated osteoblasts derived from human mesenchymal stem cells are capable of transdifferentiating into adipocytes and chondrocytes, and vice versa (11). Under osteogenic conditions, human mesenchymal stem cells increase expression of both osteogenic and adipogenic markers. Thus, osteogenic and adipogenic differentiation may constitute a set of parallel events until relatively late during osteoblast development (12). This raises the question of whether osteoporosis is a consequence of age-dependent alterations in the control of osteoblast/adipocyte fates in mesenchymal stem cells.

Unfortunately, our understanding of this control is incomplete. However, several cytokines or growth factors and associated signaling pathways, including BMP, Wnt, and polycystin pathways (13–20), are likely to play important regulatory roles. For several reasons, we hypothesized that vascular endothelial growth factor A (VEGF) may also be one of these factors. Osteoblastic precursor cells express high levels of VEGF as they migrate into the primary ossification center of developing endochondral bones in response to chemotactic factors, including VEGF (21, 22). In addition, several studies suggest that VEGF can stimulate osteoblast differentiation (23–26). Finally, VEGF expression is notably reduced in multiple cell types, including mesenchymal stem cells, with age (27–33).

To test this hypothesis, we conditionally targeted VEGF at an early stage in mesenchymal cell differentiation. We took advantage of the fact that the transcription factor osterix (Osx) (34), required for differentiation of osteoblasts, is expressed at an early stage in the osteoblast lineage. Osx- and VEGF-expressing precursor cells, located in the perichondrium of cartilage templates of endochondral bones, give rise to the majority of osteoblasts, osteocytes, and stromal cells within the primary spongiosa (trabecular bone) (22). Postnatally, mice carrying floxed alleles of VEGF and expressing Cre recombinase under the control of the Osx promoter (Osx-Cre) exhibited an osteoporosis-like phenotype characterized by reduced bone density and an increased amount of bone marrow fat. In osteoblastic and adipocytic differentiation assays with cultures of bone marrow–derived stem cells, cells from mutant animals, or cells from control animals in which VEGF was knocked down in vitro, exhibited a substantial reduction in osteoblast dif-
ferentiation and increased adipocyte differentiation. Interestingly, addition of recombinant VEGF to mutant cell cultures or addition of neutralizing antibodies against VEGF to control cells had no effects in the assays. In contrast, retrovirus-mediated restoration of VEGF expression in mutant cells increased osteoblast differentiation and reduced adipocyte differentiation to the levels exhibited by control cells. The findings suggest that VEGF produced by osteoblastic precursor cells regulates the balance of osteoblast and adipocyte differentiation via an intracrine mechanism, distinct from mechanisms involving interactions of secreted VEGF with receptors located on the surface of target cells.

Results
Mice with reduced VEGF expression in osteoblastic lineage cells exhibit reduced bone density and increased bone marrow fat. To study the skeletal functions of osteoblast-produced VEGF, we conditionally deleted Vegfa in the osteoblast lineage by crossing mice carrying floxed Vegfa alleles (Vegfafl/fl) with mice expressing Cre recombinase under the control of the Osx promoter. X-ray analysis and histological examination of 3-week-old Vegfa conditional knockout (CKO) mice showed thinner femurs and reduced trabecular bone in the proximal tibia compared with control littermates (Figure 1, A and B). These differences became more pronounced with age, and 2-month-old CKO mice showed substantially reduced bone density compared with control littermates. MicroCT analysis demonstrated a marked reduction in trabecular bone volume in mutant tibia (Figure 1C and Figure 2A) and femur (Figure 1D and Figure 2A). In femur, trabecular thickness was also reduced (data not shown). Cortical bone thickness (Figure 1, C and D) and bone volume (Figure 2, B and C) were reduced; however, since the overall bone size was smaller, cortical bone volume as percentage of total volume was not changed (Figure 2, B and C). Analysis of histological sections revealed not only decreased trabecular bone but also greatly increased adipocyte numbers in bone marrow of mutant mice (Figure 2E). Since reduced bone density could result from an imbalance between bone formation and resorption, we used tartrate-resistant acid phosphatase (TRAP) staining to assess the number of osteoclasts in bone sections. Consistent with expression of Osx-Cre in hypertrophic chondrocytes (22) and the known stimulatory effect of hypertrophic chondrocyte-generated VEGF on osteoclast recruitment into growth plate regions (35), the number of osteoclasts in the region immediately adjacent to the tibial growth plate was reduced in mutant mice compared with control littermates (Figure 2D). Thus, the reduction in bone mass was likely the result of increased bone resorption, but rather of decreased bone formation.

Histology showed no difference in trabecular bone and bone marrow fat content between Vegfafl/fl mice and heterozygous
Vegfafl/+:Osx-Cre mice (Figure 2E). Furthermore, histomorphometric analyses of the proximal tibial region showed no difference in several dynamic and static parameters between Vegfafl/fl and Osx-Cre mice (Supplemental Table 1; supplemental material available online with this article; doi:10.1172/JCI61209DS1). We therefore used Vegfafl/fl mice as control mice in the following studies. We analyzed both males and females by histology and found no sex differences regarding reduced trabecular bone density and increased bone marrow fat. To reduce variation, we used only male litter mates for all the experiments reported here. Histomorphometry of the proximal tibial region in 8-week-old Vegfafl/fl and Vegfafl/fl;Osx-Cre mice confirmed that deletion of VEGF in the osteoblast lineage resulted in a significant decrease in trabecular bone volume (Figure 3A) and trabecular number (Figure 3B) in Vegfafl/fl;Osx-Cre mice. Bone formation rate when expressed per bone surface (Figure 3D) was not affected, but bone formation rate per tissue volume was decreased (Figure 3E). Similar to bone formation rate, there was no change in osteoblast surface per bone surface and osteoblast number per bone perimeter (Supplemental Table 2). However, osteoblast number per tissue area was significantly reduced (Figure 3C), reflecting an overall decrease in osteoblast numbers relative to total bone marrow. In contrast, deletion of Vegfa resulted in 13-fold increased adipocyte numbers (Figure 3F) in the bone marrow of Vegfafl/fl;Osx-Cre mice. TRAP staining showed no change in osteoclast surface and osteoclast number within the proximal tibia secondary spongiosa (Supplemental Table 2) used for histomorphometry, although a clear trend toward decreased numbers in the mutant was apparent. These data are consistent with the possibility that the decreased trabecular bone volume observed in Vegfa CKO mice is mainly the consequence of a decrease in overall osteoblast number. In addition, the data suggest that deletion of VEGF in the osteoblast lineage stimulates differentiation of mesenchymal stem cells to adipocytes.

Reduced VEGF expression in cells of the osteoblast lineage leads to reduction in osteoblastogenesis and increased adipocyte differentiation in bone marrow cell cultures. To address the possibility that reduced osteoblastic VEGF expression may result in reduced osteoblast formation by mesenchymal stem cells in the trabecular bone marrow,
we performed colony-forming unit-fibroblast/osteoblast (CFU-F) assays with bone marrow cells from femur and tibia of mutant and control mice. In these assays, bone marrow cells are cultured at a relatively low density, allowing mesenchymal (stromal) cells to adhere and proliferate to form colonies. Each colony represents the clonal expansion of one mesenchymal cell; thus, the assay reflects the number of mesenchymal cells present in the original bone marrow cell suspension and their clonogenic potential. The number of alkaline phosphatase–positive (ALP-positive) colonies is proportional to the number of progenitors with osteoblastic potential within the mesenchymal cell population (36).

To ensure that the colonies in the CFU-F assays with bone marrow cells obtained from VEGF mutant mice were targeted by Osx-Cre;Rosa26LacZ mice. Staining of the colonies for LacZ (data not shown) revealed that the colonies were positive for LacZ, indicative of Cre activity. As shown in Figure 4, A–F, the number of ALP-positive colonies was reduced in CFU-F cultures of bone marrow cells derived from Vegfa CKO mice. When cultures were seeded with total bone marrow cells, a dramatic reduction in the total number of colonies and the fraction of colonies that were positive for ALP was seen in mutant cultures compared with controls. The total number of colonies stained with methylene blue in cultures from mutant mice was reduced to 46% of that with control cultures from littermates at day 12 (Figure 4C). Only 14% of the colonies were ALP positive in mutant cultures, compared with 27% in control littermate cultures (Figure 4B). By day 18, the total number of colonies from mutant mice stained with methylene blue was reduced to 58% of that with control littermate cultures (Figure 4F); 25% of the colonies were ALP positive in mutant cultures, while 42% were positive in control cultures (Figure 4E). Addition of recombinant VEGF to the cultures (at concentrations of 20 ng/ml or higher, see Methods) had no effect on the total number or the percentage of ALP-positive colonies (Figure 4, A–F). Furthermore, addition of neutralizing antibody against VEGF (at concentrations 5 times higher than the dose required for reducing VEGF164-induced migration of human umbilical vein endothelial cells to 50%) did not change the number of ALP-positive colonies (Figure 4, G and H). To ensure that concentrations of recombinant VEGF and neutralizing antibodies against VEGF were optimal for detecting effects of extracellular VEGF in the bone marrow cell cultures, we also assayed for osteoclast differentiation. VEGF is known to stimulate osteoclast formation in a paracrine manner (37, 38), and as expected, the number of TRAP-positive osteoclasts was significantly reduced in cultures of total mutant bone marrow cells compared with control cells (Figure 4, I and J). Addition of recombinant VEGF (20 ng/ml) to mutant cultures increased osteoclast numbers to control values (Figure 4, I and J); addition of neutralizing antibodies against VEGF to control cultures reduced osteoclast formation to the levels of mutant cell cultures (Figure 4K).

To address the possibility that the reduced number of CFU-F colonies in bone marrow cultures from mutant mice could be the result of a systemic effect in vivo prior to harvesting of the bone marrow, we isolated total bone marrow cells from control Vegfa CKO mice and knocked down VEGF expression in vitro by infection with Cre adenovirus. In CFU-F assays using these cells, results were similar to those obtained when bone marrow cells were isolated from Vegfa CKO mice (Figure 5A). Addition of recombinant VEGF to cultures treated with Cre adenovirus and of neutralizing antibody against VEGF to cultures treated with GFP adenovirus did not change the number of ALP-positive colonies (Figure 5A).

To exclude the possibility that the effects of VEGF deficiency in the CFU-F assays with total bone marrow cells may be mediated by hematopoietic cells in the bone marrow cultures, we isolated mesenchymal (stromal) cells (BMSCs) from bone marrow of control WT mice, treated the cells with lentiviral control shRNA or VEGF-specific shRNA, and performed CFU-F assays following selective growth in puromycin-containing medium. In these highly enriched BMSC cultures, 70% of the colonies were ALP positive when the cells were treated with control shRNA, and this was reduced to 33% when the cells were treated with VEGF shRNA (Figure 5, B and C). This reduction (about 50%) in the number of ALP-positive colonies was similar to the reduction in the number of ALP-positive colonies when cultures of total bone marrow cells from Vegfa/−/− and Vegfa/−/−;Osx-Cre mice were compared (Figure 4, B, C, E, and F), providing strong evidence that the CFU-F assays described above represent a robust measure of the effect of VEGF expression on the osteoblastic differentiation potential of BMSCs.

Since histology and morphometry demonstrated not only decreased trabecular bone volume but also a substantial increase
Figure 4
Suppressed osteoblastogenesis resulting from VEGF deficiency in CFU-F assays. Reduced number of ALP-positive colonies in CFU-F assays from Vegfa CKO (middle) compared with control (left) mice at day 12 (A) and day 18 (D). Staining for ALP activity using fast blue BB. Numbers of colonies with cells from mutant mice are the same in the absence and presence of exogenous VEGF (20 ng/ml) (right) in the CFU-F assay. (B and E) Comparison of colony number in the CFU-F assays shown in A and D. *P < 0.01; n = 3. (C and F) Total colonies (stained with methylene blue) in CFU-F assays at day 12 and day 18. *P < 0.001; n = 3. (G) VEGF neutralizing antibody (0.8 μg/ml) does not reduce the number of ALP-positive (fast blue BB–stained) colonies. (H) VEGF neutralizing antibody does not reduce the number of ALP-positive colonies in the assays shown in G; n = 3. (I) Osteoclastogenesis with bone marrow cells from Vegfa CKO mice (middle) is reduced compared with cells from control mice (left). Adding VEGF (20 ng/ml) to the mutant culture restores osteoclastogenesis to control levels (right). Original magnification, ×10. (J) Quantitation of osteoclastogenesis data as presented in I. (K) Osteoclastogenesis with bone marrow cells from control mice is reduced when neutralizing VEGF antibodies (0.8 μg/ml) are added to the culture. Original magnification, ×10.
Figure 5
Loss of VEGF in osteoblast lineage increases adipogenesis. (A) Reduced numbers of ALP-positive colonies following addition of Ad-Cre to bone marrow cells from Vegfafl/fl mice compared with control Ad-GFP. The number of colonies is not affected by exogenous VEGF (20 ng/ml) or neutralizing antibody (0.8 μg/ml). \( *P < 0.01; n = 3 \). (B) Reduced number of ALP-positive colonies in bone marrow cultures expressing VEGF shRNA compared with control shRNA. \( *P < 0.05; n = 3 \). (C) The total number of colonies is not affected by treatment of cultures with VEGF-specific shRNA; \( n = 3 \). (D) Adipocyte colonies (boundaries indicated by dashed lines) in assays used for colony counts. Original magnification, \( \times 10 \). (E) More adipocyte colonies in CFU-A assays with BMSCs of mutant mice (middle) in adipogenic medium than with cells from control littermates (left). Addition of 20 ng/ml VEGF has no significant effect. \( *P < 0.01 \) compared with Vegfafl/fl control; \( n = 3 \). (F) No effect of neutralizing antibody (0.8 μg/ml) against VEGF in CFU-A assays with cells from control mice. (G) Increased number of adipocyte colonies in assays with bone marrow cells from Vegfafl/fl mice treated with Ad-Cre compared with Ad-GFP. No effect of exogenous VEGF (20 ng/ml) or neutralizing antibody (0.8 μg/ml). \( *P < 0.01; n = 3 \). (H) Increased number of adipogenic colonies in cultures expressing VEGF shRNA compared with control shRNA. \( *P < 0.001; n = 4 \).

Mechanisms by which VEGF stimulates osteoblast and represses adipocyte differentiation. Our finding that exogenous VEGF or neutralizing antibody against VEGF affects neither the number of ALP-positive CFU-F colonies nor the number of adipocyte colonies in CFU-A cultures raises the possibility that VEGF controls the balance between osteoblastogenesis and adipogenesis through an intracrine mechanism. If this is correct, retrovirus-mediated expression of a Vegfa gene in mutant BMSCs should rescue reduced osteoblastogenesis in CFU-F and increased adipogenesis in CFU-A assays. When we infected BMSCs — isolated from Vegfafl/fl mice and treated with Cre adenovirus — with VEGF retrovirus (Re-VEGF) or control GFP retrovirus (Re-GFP), the retrovirus-mediated expression of VEGF rescued the defects in CFU-F and CFU-A cultures due to Cre adenovirus–induced loss of VEGF expression (Figure 6, A and B).

These data suggest that osteoblast-generated VEGF regulates the number of CFU-F and CFU-A colonies in bone marrow cell cultures via intracrine mechanisms that do not involve extracellular VEGF. To address the question of whether these mechanisms include interaction of VEGF with its receptors, we first used the Osx-Cre strategy to conditionally target Flt1 and Flk1 in osteoblastic lineage cells. Histological examination of mutant skeletons revealed decreased trabecular bone in both mutants, but no increase in bone marrow adipocyte numbers (data not shown). As expected, based on the in vivo bone phenotypes, examination of ALP-positive CFU-F colonies in total bone marrow cell cultures from both Flt1 and Flk1 CKO mice showed that the number of positive colonies was reduced when compared with cultures from control littermates (Figure 6, C and D). In contrast, knocking down Flk1 expression had no effect on the number of adipocyte colonies in CFU-A assays and knocking down Flt1 reduced the number of adipocyte colonies compared with controls (Figure 6, E and F). Thus, both receptors are required for the stimulatory effect of VEGF on osteoblastogenesis, but they are not critical for the ability of VEGF to repress adipocyte differentiation.

in the number of adipocytes in bone marrow of Vegfa CKO mice, we next addressed the cellular mechanism of increased adipocyte formation. In vitro colony forming unit–adipocyte (CFU-A) assays showed that mutant BMSCs in adipogenic medium formed more adipocyte colonies than cells from control littermates (Figure 5, D and E). Addition of exogenous VEGF or neutralizing antibody against VEGF had no effect on the number of adipocyte colonies in these cultures (Figure 5, E and F), indicating that VEGF-dependent mechanisms underlying differentiation of adipocytes from BMSCs are not affected by extracellular VEGF. To exclude the possibility that increased adipogenesis in mutant mice is caused by a systemic effect in vivo prior to harvesting the bone marrow, we isolated BMSCs from control mice and knocked down VEGF levels in vitro using Cre adenovirus. When we analyzed adipocyte formation using these cells, results were similar to those obtained with BMSC cultures from mutant mice (Figure 5G). Finally, we treated BMSCs with lentivirus encoding control shRNA or VEGF-specific shRNA, selected the cells in puromycin-containing medium, and performed CFU-A assays in adipogenic medium. In these highly enriched BMSC cultures, the number of adipocyte colonies was significantly increased when the cells expressed VEGF-specific shRNA (Figure 5H).
Next, we stained BMSCs with antibodies against VEGF, VEGFR1, and VEGFR2 to determine whether the 3 proteins are colocalized within particular intracellular compartments. As shown in Figure 7A, positive staining for VEGF and VEGFR2 was detected in both nuclear and cytoplasmic regions, but staining for VEGFR1 was primarily observed in the nuclear area, as demonstrated by the similarity to the staining pattern seen when BMSCs were immunostained for the nuclear envelope protein lamin A/C. Based on this observation and our finding that osteoblasts isolated from femur and tibia of Vegfa<sup>fl/fl</sup>;Osx-Cre mice exhibit the flattened phenotype of senescent cells and stain positively for senescence-associated β-galactosidase (data not shown), we considered a potential association between lamin A/C and intracrine VEGF signaling. Depletion of lamin A/C induces cellular senescence (39), and knockdown of lamin A/C in BMSCs has been reported to reduce osteoblastic differentiation, increase adipocyte differentiation, and lower RUNX2 DNA binding activity (40–44). Furthermore, lamin A–null mice exhibit a postnatal decline in bone mass (44). Given the similarity of these characteristics to those of mice lacking VEGF in osteoblastic lineage cells, we considered the possibility that the effect of VEGF on osteoblastogenesis may be mediated by a mechanism that includes a functional interaction between VEGF and lamin A/C.

First, we used quantitative assays for VEGF and Western blotting for lamin A/C to analyze cell lysates obtained from BMSCs of control (WT) and heterozygous lamin A/C–null (Lmna<sup>−/−</sup>) mice treated with lentiviral control shRNA or VEGF-specific shRNA. As shown in Figure 7B, the concentration of cell-associated VEGF was reduced to almost the same extent in lysates obtained from cells expressing VEGF-specific shRNA and control shRNA-expressing cells from Lmna<sup>−/−</sup> mice. In addition, Western blotting demonstrated that protein levels of lamin A (Figure 7C) were increased in cells expressing VEGF-specific shRNA. These data suggest that while lamin A stimulates VEGF protein expression, VEGF represses levels of lamin A. When the lentivirus-treated cells were used for CFU-F assays, we found that Lmna<sup>−/−</sup> cells generated more total colonies than WT cells (Figure 7E), consistent with published evidence that lamin A/C represses proliferation of mesenchymal cells (45–47). However, the fraction of ALP-positive colonies in the assays was proportional to the level of VEGF protein in the cells (Figure 7, B, D, and E). These data indicate that cell-associated VEGF is a major stimulator of differentiation of BMSCs along the osteoblastic pathway. In contrast, cell-associated VEGF represses differentiation of BMSCs to adipocytes. When CFU-A assays were carried out using lentivirus-treated BMSCs from WT and Lmna<sup>−/−</sup> mice, the relative number of adipocyte colonies increased dramatically when the levels of VEGF in cell lysates at the start of the CFU-A assay were low (Figure 7F). However, the VEGF levels in the culture medium were hardly affected when cells expressed VEGF-specific shRNA and showed no correlation with adipocyte colony numbers (Figure 7G). The number of adipocyte colonies as a fraction of total colonies (about 30%) was similarly increased in WT cultures express-
Figure 7
Mechanisms by which VEGF controls osteoblast/adipocyte differentiation in BMSCs. (A) VEGF and VEGFR2 expression is detected in both nuclear and cytoplasmic regions of Osx-expressing cells, whereas VEGFR1 expression is primarily observed in the nuclear area similar to lamin A/C. Original magnification, ×250. (B) Reduced VEGF protein levels associated with WT and Lmna+/– cultures expressing VEGF-specific shRNA. *P < 0.05, **P < 0.01; n = 3. (C) Increased lamin A protein levels associated with WT and Lmna+/– cultures expressing VEGF-specific shRNA. Values above and below the blots represent relative lamin A and β-actin protein levels. (D) Reduced number of ALP-positive colonies in both WT and Lmna+/– cultures expressing VEGF shRNA. *P < 0.05, **P < 0.01; n = 3. (E) The number of colonies increased in Lmna+/– compared with WT cultures but was not affected by treatment with VEGF-specific shRNA. *P < 0.05; n = 3. (F) Increased number of adipogenic colonies in WT and Lmna+/– cultures expressing VEGF shRNA and Lmna+/– cultures expressing control shRNA. *P < 0.001; n = 4. (G) VEGF protein levels in culture medium prior to CFU-A assay are only slightly affected in WT and Lmna+/– cultures expressing VEGF-specific shRNA. *P < 0.05 and **P < 0.01. (H) VEGFR1 is mainly expressed as the soluble splice variant (sVEGFR1), whereas full-length (I) and phosphorylated (J) VEGFR2 is detected in WT and Lmna+/– cultures. (K) Reduced RUNX2 transcriptional activity in cultures expressing VEGF-specific shRNA and/or Lmna+/– cultures expressing control shRNA. *P < 0.05; n = 4. (L and N) RUNX2 protein levels are decreased in WT (L) and Lmna+/– (N) cultures expressing VEGF-specific shRNA under osteogenic conditions. (M and O) PPARγ2 protein levels are increased in WT (M) and Lmna+/– (O) cultures expressing VEGF-specific shRNA in the presence of adipogenic inducers.

Discussion
In this study, VEGF was conditionally targeted in Osx-positive osteoblast lineage cells, allowing studies of the functions of osteoblast-derived VEGF. Our data demonstrate that osteoblast-derived VEGF stimulates formation of osteoblasts and suppresses adipogenesis, both in vivo and in vitro. Thus, normal postnatal bone homeostasis requires expression of VEGF in osteoblasts. The reduction in bone mass and the increase in marrow fat in Vegfa CKO mice is similar to changes associated with osteoporosis and age-related osteopenia (4). Postmenopausal women with polymorphisms associated with high or low VEGF production have higher or lower lumbar spine bone mineral density, respectively (49). Intriguingly, there also appears to be no significant association between levels of circulating VEGF and bone mineral density (49). This is consistent with our in vitro data demonstrating that addition of recombinant VEGF to cultures of VEGF-deficient BMSCs, even at very high concentrations, has no effects on the differentiation properties of the mutant cells. This suggests that VEGF controls osteoblastogenesis and adipogenesis via mechanisms that are resistant to the effects of extracellular VEGF.

The concept of an intracellular signaling loop for VEGF has been previously suggested in the case of bone marrow hematopoietic stem cells (50). VEGF appears to control survival of these stem cells via a mechanism that is resistant to inhibitors that fail to penetrate the plasma membrane, such as antibodies (50). A similar phenomenon has been noticed in studies in which VEGF was specifically deleted in endothelial cells (51). Furthermore, a recent report suggests that VEGF increases survival and chemoresistance in human colorectal cancer cells by an intracellular mechanism (52). How intracellular functions of VEGF may be regulated and in which compartments VEGF may function intracellularly has not been known. However, several observations provide intriguing hints. VEGF has been reported to mediate survival of human breast cancer carcinoma cells via binding intracellularly to VEGFR1 (53), and it has been found to accumulate in endothelial cell nuclei (54). The immunocytochemical detection of sVEGFR1, VEGFR2, and VEGF in the nuclear region of BMSCs in the present study is consistent with these observations. A key question is whether the intracellular function in Osx-positive osteoblastic lineage cells requires direct binding of VEGF to its receptors. Our data do not allow a definitive answer to this question. However, since knockdown of either Flk1 or Fkh1 in the osteoblast lineage led to reductions in the numbers of ALP-positive CFU-F colonies, interactions of intracellular VEGF with sVEGFR1 and VEGFR2 in differentiating osteoblasts may be part of the mechanism by which VEGF stimulates osteoblastic differentiation in BMSCs. In lysates of BMSCs, VEGFR2 was found to be phosphorylated on Y1175, and downstream targets, such as Akt and p38MAPK, were phosphorylated as well even when cell-associated VEGF protein levels were knocked down (Supplemental Figure 1). Therefore, while the receptors may be involved in the intracellular effects of VEGF on osteoblastic differentiation, it is unlikely that the effects are mediated by signaling downstream of activated VEGFR2. Interestingly, Domingues et al. (55) recently reported that a VEGF/VEGFR2 complex can be transported into nuclei in endothelial cells, where it can interact with several nuclear proteins, including the transcription factor Sp1, and stimulate the expression of VEGFR2. If this mechanism also applies to BMSCs, one would expect the level of VEGFR2 to decrease with decreasing levels of VEGF in the cells, and this is consistent with lower levels of VEGFR2 in lysates from BMSCs treated with VEGF-specific shRNA.
The association between low levels of VEGF and low levels of RUNX2 protein and activity in BMSCs provides an explanation for the reduced ability of the cells to differentiate into osteoblasts when VEGF levels are knocked down. Elucidation of the mechanisms by which VEGF affects RUNX2 expression and activity will require further studies. However, since Sp1 has been demonstrated to be important for stimulation of Runx2 transcription (36), the findings of Domingues et al. (55) raise the possibility that VEGF/VEGFR2 complexes may have a role in Sp1-dependent induction of Runx2 transcription. Combined with evidence that RUNX2 can serve as a transcription factor to induce transcription of Vegfa (35, 57), this suggests the existence of a positive feedback loop between intracrine VEGF and RUNX2 expression.

The suppression of adipocyte differentiation by VEGF does not include VEGFR1- and VEGFR2-dependent signaling mechanisms. Results of CFU-A assays with VEGF-R2-deficient BMSCs were not different from those of assays with control cells. Also, VEGFR1 deficiency in BMSCs inhibited adipogenesis. This is consistent with the known function of sVEGFR1 as a high-affinity decoy receptor for VEGF in many contexts; loss of VEGFR1 may increase the efficiency of VEGF as a suppressor of adipogenesis. Thus, other VEGF-binding receptors or receptor-independent VEGF functions may be involved in the mechanism by which VEGF suppresses adipocyte differentiation. Knockdown of lamin A in mesenchymal stem cells results in increased adipocyte differentiation (42, 43), raising the question of whether the effect of VEGF on adipocyte differentiation is mediated by lamin A. Our data indicate that this is not the case. Loss of one Lmna allele in BMSCs caused cell-associated VEGF to be reduced to the levels in cells treated with VEGF-specific shRNA, suggesting that lamin A stimulates VEGF protein expression. In addition, the fact that levels of lamin A/C expression increased when WT or Lmna°/° mesenchymal cells expressed VEGF-specific shRNA indicates that VEGF, in turn, suppresses lamin A expression. Such a reciprocal relationship between VEGF and lamin A may explain why loss of one Lmna allele in cells results in lamin A levels that are greater than 50% of what they are in WT cells. The relationship between VEGF and lamin A also indicates that lamin A is upstream of VEGF with respect to both osteoblast and adipocyte differentiation. In fact, our data suggest that most, if not all, effects of lamin A on osteoblast and adipocyte differentiation may be mediated by the effects of lamin A on the level of VEGF in cells (Figure 8).

Activation of Wnt/β-catenin signaling suppresses adipogenesis and promotes osteoblastogenesis of mesenchymal stem cells. This raises the question of whether Wnt signaling may be involved in the VEGF-dependent functions in the cells (13–15). However, disruption of canonical Wnt signaling in mice causes bone loss due to reduced osteoblast differentiation, but adipogenesis is not affected (16). Furthermore, transdifferentiation of mesenchymal stem cell-derived osteoblast progenitors into adipocytes is not associated with a change in expression of Wnt3a and β-catenin (58).

In addition to the intracrine mechanism for VEGF function related to osteoblast/adipocyte differentiation as suggested by the data presented here, we also demonstrate that VEGF secreted by BMSCs has paracrine effects on osteoclast differentiation in assays using total bone marrow cell cultures. Furthermore, it is known that osteoblast-derived VEGF can affect bone formation in a paracrine manner via stimulation of angiogenesis (59). It will be interesting to investigate whether and how osteoblast-derived intracrine VEGF affects the interaction between osteoblasts and endothelial cells in bone marrow, since it has been reported that direct contact between endothelial cells and osteoprogenitors in vitro supports osteoblast function (60). To our knowledge, this is the first in vitro/in vitro study to investigate the roles of osteoblast-derived VEGF in osteoblastic and adipocytic differentiation. Based on the results of the study, we propose that understanding the mechanisms by which intracrine VEGF controls adipocyte and osteoblast fates may lead to identification of novel therapeutic targets to control bone loss in osteoporosis.

**Methods**

*Mice strains.* Vegfa floxed and Ossx-Cre mouse lines have been described previously (61, 62). Flk1 floxed and Flk1 floxed mice were generated using standard procedures. Genomic DNA isolated from portions of mouse tails were used for VEGF and VEGF receptor genotyping. Genotyping for Vegfa was carried out using PCR with the following primers: forward 5′-CCTG-GCCCTCAAGTACACCTT-3′; reverse 5′-TCCGTAGCACGGCATTTCTC- TAG-3′. For Flk1 the primers were forward 5′-CCTGCTAGTATTCCCT- GATTGGA-3′; reverse 5′-GCCCATAGCTACCTGGC-3′. For Flk1 the primers were forward 5′-GACTTGGTTCTATCAGCCTAG-3′; reverse 5′-GACGCTTGTAAGCTGCTACAC-3′. The primers for Cre were forward 5′-GATGAGGTTCGCAAGAACCTG- GTGCAAATC-3′; reverse 5′-TGAACGAACCTGGTCTGAAAATC-3′. The Rosa26 LacZ reporter mice and the Lmna°/° mice were obtained from the Jackson Laboratory. To assess the efficiency of Ossx-Cre-mediated inactivation of Vegfa in bones of mutant mice, RNA was extracted from tibia and femur of newborn control and mutant pups and used for real-time PCR with primers (forward 5′-ATCTTCAAGCC-GTCCCTGTG-3′ and reverse 5′-CTGATGTTGATGTGCT-3′) from within exon 3 (flanked by loxP sites in the Vegfa gene). Real-time PCR for Gapdh transcripts (forward primer 5′-CCTGCTACGACGCCACCT-3′ and reverse primer 5′-GCCTAAGCTCACCTGG-3′) served as internal control. The results indicated that exon 3−containing Vegfa transcripts in extracts of bones were reduced by about 70% in mutant bones. Since isolated bones contain non-osteoblastic cells, such as endothelial cells, this value likely underestimates the degree of Ossx-Cre−dependent Vegfa deletion in osteoblastic cells.

**X-ray and microCT.** X-ray and microCT analyses were performed using Faxitron Micro 50 X Ray and Scanco Medical μCT-40 instruments.

**Histology and histochemistry.** Mouse limbs were fixed in 4% paraformaldehyde and embedded in paraffin for standard histology or dehydrated with 20% sucrose and embedded in Tissue-Tek O.C.T. compound (Sakura) for cryostat sectioning. For lipid staining, frozen sections or cell cultures were fixed with 10% neutral buffered formalin and stained with filtered oil red O in 60% isopropanol. ALP staining of cell cultures was performed using Alkaline Phosphatase kit 86R-1KT (with fast red violet LB) or 86C-1KT (with fast blue BB) (Sigma-Aldrich) as described in the manufacturer’s instructions. TRAP staining of frozen sections for osteoclasts was performed using Alkaline Phosphatase kit 86R-1KT (with fast red violet LB) or 86C-1KT (with fast blue BB) (Sigma-Aldrich) as described in the manufacturer’s instructions.
sides and cultured in 8- to 12-week-old mice were flushed out with ice-cold α
clature (63). A sampling site with an area of approximately 1.2 mm² was
number was counted using toluidine blue–stained sections. Histomorpho-
quantify osteoblast number and osteoclast number, respectively. Adipocyte
TV]). Consecutive sections were stained with toluidine blue and TRAP to
surface [BFR/BS], bone volume [BFR/BV], and tissue volume referent [BFR/
mineralizing surface (MS/BS) and bone formation rate expressed per bone
Cre (6 mice) mice were subcutaneously injected with 20 mg/kg calcein and
mice for each genotype were plated in triplicate dishes. 

scored by counting clusters containing more than 6 cells in the entire dish. 
were cultured for 3–4 days before oil red O staining and counting adipo-
M 1-isobutyl-3-methylxantine (IBMX) were added, and cells

Isolation of osteoblastic cells from mouse long bones. Isolation and culture of 
primary bone cells from tibia and femur of 8- to 12-week-old mice were as 
described (64). Bone marrow cells were flushed out with PBS, and cleaned 
diaphyses were cut into small pieces and incubated in bacterial collage-
nase solution at 37°C. The bone pieces were cultured in DMEM contain-
ing 15% FBS and 100 µg/ml ascorbic acid until cells migrating from the 
bone chips became confluent.

Isolation of bone marrow cells. Bone marrow cells from tibia and femur of 
8- to 12-week-old mice were flushed out with ice-cold α-MEM with nucleo-
sides and cultured in α-MEM containing 10% or 20% FBS following lysis of 
red blood cells with RBC Lysis Buffer (eBioscience).

For CFU-F assays, 1 × 10⁶ bone marrow cells were plated on 60-mm dishes 
in 5 ml α-MEM (Invitrogen) containing penicillin/streptomycin, 50 µg/ml 
ascorbic acid (Sigma-Aldrich), 10 µM dexamethasone (Sigma-Aldrich), and 
10% FBS (Atlanta Biologicals). After 12 or 18 days, cultures were fixed with 
70% ethanol and stained for ALP (using fast red violet LB or fast blue BB) 
or with methylene blue for counting of all colonies. CFU-F colonies were 
formed with 10% horse serum, 10–8 M hydrocortisone, 60 

GFP adenovirus (Ad-GFP) was used as control.

Isolation of osteoblastic cells from mouse long bones. Isolation and culture of 
primary bone cells from tibia and femur of 8- to 12-week-old mice were as 
described (64). Bone marrow cells were flushed out with PBS, and cleaned 
diaphyses were cut into small pieces and incubated in bacterial collage-
nase solution at 37°C. The bone pieces were cultured in DMEM contain-
ing 15% FBS and 100 µg/ml ascorbic acid until cells migrating from the 
bone chips became confluent.

For CFU-F assays, 1 × 10⁶ bone marrow cells were plated on 60-mm dishes 
in 5 ml α-MEM (Invitrogen) containing penicillin/streptomycin, 50 µg/ml 
ascorbic acid (Sigma-Aldrich), 10 µM dexamethasone (Sigma-Aldrich), and 
10% FBS (Atlanta Biologicals). After 12 or 18 days, cultures were fixed with 
70% ethanol and stained for ALP (using fast red violet LB or fast blue BB) 
or with methylene blue for counting of all colonies. CFU-F colonies were

Isolated cells were cultured in α-MEM containing 10% or 20% FBS following lysis of red blood cells with RBC Lysis Buffer (eBioscience).

For CFU-F assays, 1 × 10⁶ bone marrow cells were plated on 60-mm dishes in 5 ml α-MEM (Invitrogen) containing penicillin/streptomycin, 50 µg/ml ascorbic acid (Sigma-Aldrich), 10 µM dexamethasone (Sigma-Aldrich), and 10% FBS (Atlanta Biologicals). After 12 or 18 days, cultures were fixed with 70% ethanol and stained for ALP (using fast red violet LB or fast blue BB) or with methylene blue for counting of all colonies. CFU-F colonies were scored by counting clusters containing more than 6 cells in the entire dish. For each experiment, repeated at least 3 times, bone marrow cells from 2–3 mice for each genotype were plated in triplicate dishes.

For CFU-A assays, bone marrow cells were plated at 2 × 10⁶ cells per well in 24-well plates with α-MEM containing 20% FBS. When cells became 70%–80% confluent, adipocyte differentiation medium (DMEM supplemented with 10% horse serum, 10 µM hydrocortisone, 60 µM indomethacin, and 500 µM 1-isobutyl-3-methylxantine (IBMX) were added, and cells were cultured for 3–4 days before oil red O staining and counting adipocyte clusters in each well. Each experiment (more than 3 were performed) included triplicate or quadruplicate wells.

In vitro experiments with BMSCs. Bone marrow cells from tibia and femur of 8- to 12-week-old mice were cultured in α-MEM containing 10% FBS. Non-adherent cells were removed by replacing the medium after 3 days. The attached BMSCs were used for experiments. Cells from Vegfaβ mice were treated with Cre adenovirus (Ad-Cre) in vitro to knock down Vegfa expression. GFP adenovirus (Ad-GFP) was used as control.

For experiments requiring retroviral infection, pWZL Blast VEGF and pWZL Blast GFP plasmids, generated by B. Weinberg, were provided by Addgene (cat. 10909 and 12269). The plasmids were transfected into 293-GPG packaging cells using FuGENE 6 (Roche). After 48 hours, superna-
tants were collected and used for infection. The BMSCs (isolated from 
Vegfaβ mice and treated with Ad-GFP or Ad-Cre to knock down Vegfa) were infected with VEGF retrovirus (Re-VEGF) or GFP retrovirus (Re-GFP).

For lentiviral shRNA experiments, bone marrow cells from WT and 
Lmna–/– mice were isolated and treated with VEGF shRNA Lentiviral Par-
ticles (sc-36815-V, Santa Cruz Biotechnology Inc.), Control shRNA Lentivi-
Rental Particles (sc-108080, Santa Cruz Biotechnology Inc.), or cGFP Control 
Lentiviral Particles (sc-108084, Santa Cruz Biotechnology Inc.) in the pres-
ence of 5 µg/ml Polybrene (sc-134220, Santa Cruz Biotechnology Inc.). To 
select stable clones expressing the shRNA, cells were treated with 2 µg/ml puromycin dihydrochloride (sc-108071, Santa Cruz Biotechnology Inc.) until resistant colonies were identified and used for subsequent experiments.

ELISA assays. VEGF protein levels in cell lysates were assessed using the 
Quantikine Mouse VEGF Immunoassay (R&D Systems) in accordance 
with the manufacturer’s instructions; VEGF levels in lysates from confluent cultures were normalized to protein levels. For measurements of VEGF in culture media, media were collected when cells were 70%–80% confluent and ready to be grown in adipogenic medium for 3 days for CFU-A assays.

The optical density of each well was determined using a microplate reader set to 450 nm. The optical density determined with blank culture medium was used as a control.

Immunocytochemistry. Cells seeded in culture chambers were fixed in 4% PFA and incubated with antibodies against VEGF (sc-152, Santa Cruz Biotech-

nology Inc.), Flt-1 (sc-316, Santa Cruz Biotechnology Inc.), Flk-1 (sc-505, 
Santa Cruz Biotechnology Inc.), or lamin A/C (sc-20681, Santa Cruz Bio-

technology Inc.) in combination with OSX (sc-22538, Santa Cruz Bio-
technology Inc.) antibodies at 1:50 dilution for 60 minutes at room tem-

perature. After 3 washes with PBS, cells were incubated with Alexa Fluor 
555–labeled donkey anti-rabbit IgG and Alexa Fluor 488–labeled donkey anti-goat IgG (Invitrogen) at 1:200 dilution for 45 minutes. DAPI was used for nuclear staining. Fluorescence images were obtained with a Nikon 80i Upright microscope and associated software.

Western blotting. Cell lysates were prepared by adding ice-cold Phospho-
Safe Extraction Reagent (Novagen) supplemented with protease inhibitor 
cocktail (Roche). After centrifugation for 20 minutes (16,500 g) at 4°C, 
protein concentrations of supernatants were determined by Bradford Pro-
tein Assay kit (Thermo Scientific). Equal amounts of protein were loaded 
and separated by SDS-PAGE using 4%–12% Bis-Tris precast polyacrylamide 
gels (NUPAGE; Invitrogen), followed by transfer of proteins onto nitrocel-
ulose membranes (Bio-Rad). Blots were blocked with 5% Blotting-Grade 
Blocker Nonfat Dry Milk (Bio-Rad) in TBS/0.1% (v/v) Tween-20, followed 
by incubation overnight with primary antibodies against lamin A/C (1:500, 
sc-20681, Santa Cruz Biotechnology Inc.), Flk-1 (1:500, sc-316, Santa Cruz 
Biotechnology Inc.), Flt-1 (1:500, sc-505, Santa Cruz Biotechnology Inc.), 
phospho-VEGFR2 (1:1,000, 2478, Cell Signaling Technology), RUNX2 
(1:1,000, 8486, Cell Signaling Technology), PPARγ (1:1,000, 2443, Cell Sig-
naling Technology), phospho-Akt (Ser473, 1:1,000, 9271, Cell Signaling 
Technology), phospho-p38 MAP Kinase (Thr180/Tyr182, 1:1,000, 9211, 
Cell Signaling Technology), and β-actin (1:5,000, 5441, Sigma-Aldrich) in 
5% BSA in TBS/0.1% (v/v) Tween-20. Following incubation with horseradish 
peroxidase–conjugated secondary antibodies for 1 hour, immunoreactive 
bands were detected by chemiluminescent substrate (Thermo Scientific).

DNA transfections and reporter assays. Bone marrow cells were plated at 
2.5 × 10⁴ cells per well in 24-well plates and transfected with 500 ng OSE-
luc (a gift from Toshihisa Komori, Nagasaki University Graduate School 
of Biomedical Sciences, Nagasaki, Japan) and 12.5 ng of a thymidine 
kinase promoter–driven Renilla luciferase construct (Promega) per well, 
using Lipofectamine LTX Plus Reagent (Invitrogen). After transfection, 
cells were cultured for 24 hours in osteogenic culture medium containing 
100 µM ascorbic acid 2-phosphate (Wako) and 8 mM β-glycerophosphate
(Sigma-Aldrich). Cells were then lysed, and extracts were used in luciferease assays performed with the Dual-Glo Luciferase Assay System (Promega) in accordance with the manufacturer’s instructions. Luciferase activity was measured using a microplate luminesimeter, and the firefly luciferase value in each well was normalized to the corresponding Renilla luciferase readout.

Blocking antibodies and recombinant proteins. Antibodies against mouse VEGF (affinity-purified polyclonal goat IgG, cat. AF-493-NA) from R&D Systems were used for blocking experiments. The antibody concentration (0.8 μg/ml) in the blocking experiments with anti-VEGF is the upper limit for ELISA capture with this antibody and about 5 times higher than the upper range for the neutralization dose (ND50) for this antibody (0.05–0.15 μg/ml in the presence of 10 ng/ml recombinant VEGF164, measured by its ability to neutralize VEGF164-induced proliferation in human umbilical vein endothelial cells (65). This antibody concentration was sufficient to reduce osteoclast differentiation to the low levels seen with bone marrow cultures from Vegfa−/−Oxst-Cre mice when antibody was added to cultures from Vegfa−/− mice. To test whether exogenous VEGF could rescue the changes in CFU-F assays following knockdown of Vegfa in bone marrow cells, recombinant VEGF was added at concentrations of 20 ng/ml, 50 ng/ml, and 100 ng/ml. At these concentrations no rescue effects were seen. However, a concentration of 20 ng/ml of VEGF was sufficient to restore osteoclastogenesis to control levels when the protein was added to cultures of bone marrow cells from Vegfa−/−Oxst-Cre mice. Recombinant mouse VEGF (VEGF164) (cat. 493-MV/CF) was obtained from R&D Systems.

Statistics. Results are presented as mean ± SD, and unequal t-tested Student’s t-test was used. P values less than 0.05 were considered significant.

22. Swift ME, Kleinman HK, DiPietro LA. Impaired wound repair and delayed angiogenesis in aged...