PPARγ insufficiency enhances osteogenesis through osteoblast formation from bone marrow progenitors

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Based on the fact that aging is associated with a reciprocal decrease of osteogenesis and an increase of adipogenesis in bone marrow and that osteoblasts and adipocytes share a common progenitor, this study investigated the role of PPARγ, a key regulator of adipocyte differentiation, in bone metabolism. Homozygous PPARγ-deficient ES cells failed to differentiate into adipocytes, but spontaneously differentiated into osteoblasts, and these were restored by reintroduction of the PPARγ gene. Heterozygous PPARγ-deficient mice exhibited high bone mass with increased osteoblastogenesis, but normal osteoblast and osteoclast functions, and this effect was not mediated by insulin or leptin. The osteogenic effect of PPARγ haploinsufficiency became prominent with aging but was not changed upon ovariectomy. The PPARγ haploinsufficiency was confirmed to enhance osteoblastogenesis in the bone marrow cell culture but did not affect the cultures of differentiated osteoblasts or osteoclast lineage cells. This study demonstrates a PPARγ-dependent regulation of bone metabolism in vivo, in that PPARγ insufficiency increases bone mass by stimulating osteoblastogenesis from bone marrow progenitors.

Introduction

Osteoblasts and adipocytes share a common progenitor: multipotential mesenchymal stem cells in bone marrow (1–3). Accumulated evidence of the differentiation switching of these two cell lineages suggests that a large degree of plasticity exists between them and that the relationship is reciprocal (4–6). The clinical fact that a decrease in bone volume (BV) of age-related osteoporosis is accompanied by an increase in marrow adipose tissue (7–9) also implies the possible reciprocal relationship that is postulated to exist between the two differentiation pathways. The signal transduction pathways implicated in this process are therefore evaluated as potential targets for therapeutic intervention of osteoporosis. The molecular mechanism underlying the reciprocal relationship is not yet well understood, however, although several studies using strain-specific and KO murine models have begun to explore the relationship in vivo (10–15).

Several key transcription factors that function in the complex transcriptional cascade during adipocyte differentiation have been identified, including PPARγ and CCAAT enhancer-binding proteins (C/EBPs) (16). PPARγ is a ligand-activated transcription factor that belongs to the nuclear hormone receptor superfamily and functions as a heterodimer with a retinoid X receptor by binding to the PPAR responsive element (PPRE) within the promoters of the target genes (17–19). PPARγ is expressed early in the adipocyte differentiation program and is activated by long-chain fatty acids, peroxisome proliferators, and the thiazolidinedione class of antidiabetic agents (17–19). Most importantly, PPARγ plays requisite and sufficient roles in the regulation of adipocyte differentiation, because its overexpression in fibroblast cell lines initiates adipogenesis (20) and ES cells and embryonic fibroblastic cells from mice lacking PPARγ were unable to differentiate into adipocytes (21–23).

When one takes the results of the studies together, it is possible that PPARγ may contribute not only to adipogenesis, but also to osteogenesis in the bone marrow where bipotential precursors can differentiate to either adipocytes or osteoblasts. This study investigated the physiological role of PPARγ on the marrow cells and bone cells using in vivo morphological analyses and ex vivo cell culture systems. For the in vivo analysis, we used mice lacking the PPARγ gene, which we generated by gene targeting (22). Although the homozygous PPARγ-deficient (PPARγ−/−) mice were embryonically lethal at 10.5–11.5 days after post coitum due to placental dysfunction, heterozygous PPARγ-deficient (PPARγ+/−) mice developed normally. The heterozygotes led to a 50% reduction in PPARγ expression and exhibited resistance to high-fat diet–induced obesity and insulin resistance; however, on a standard diet they grew normally, without abnormalities in major organs such as brain, heart, liver, spleen, or kidney (22, 24). We show here that the homozygous PPARγ-deficient ES cells spontaneously differentiate into osteoblasts ex vivo and that PPARγ haploinsufficiency due to the heterozygous PPARγ deficiency resulted in enhanced bone formation with increased osteoblastogenesis from bone marrow progenitors both in vivo and ex vivo.

Methods

Animals. The generation of PPARγ gene–targeted mice was described previously (22). In each experiment, WT and PPARγ+/−
mice littermates fed a standard diet were compared. All experiments were performed on male mice at 8 or 52 weeks of age, except for the ovariectomy (OVX) experiment in which female mice underwent surgical operation at 26 weeks and were analyzed at 30 weeks. All experiments were performed according to the protocol approved by the Animal Care and Use Committee of the University of Tokyo.

ES cell cultures. Mouse PPARγ–/– and WT ES cells were isolated from blastocysts generated by mating PPARγ–/– mice with each other, as previously described (25). ES cells were maintained in DMEM supplemented with 15% FBS, 200 mM L-glutamine, 100 µM β-mercaptoethanol, and 104 U/ml of leukemia inhibitory factor (LIF; Chemicon International, Temecula, California, USA). Differentiation of ES cells was induced by using a modified protocol, described previously (2, 26). In brief, after being trypsinized with 0.025% trypsin-EDTA, cells were plated onto a bacterial Petri dish in the absence of LIF and cultured with 100 nM all-trans retinoic acid for 5 days, with medium being replenished on day 3. The embryoid bodies were transferred to a gelatinized six-multiwell plate and allowed to adhere to the well with DMEM containing 10% FBS. For the analysis of osteogenic differentiation, cultures were maintained in the same condition without any additional supplements for 10 days, were fixed with 10% buffered formalin, and were incubated in the presence of 5% silver nitrate solution under an ultraviolet light for 10 minutes, then incubated for 5 minutes in the presence of 5% sodium thiosulfate solution (von Kossa staining). To discern the calcified nodules from the embryoid body, both of which are seen as black, the von Kossa–positive nodules that do not connect to the embryoid body in a well were counted. For the analysis of adipogenesis, the medium was supplemented with 1 µM troglitazone (Sankyo Pharmaceutical Co., Tokyo, Japan) for 10 days, fixed in 10 mM sodium periodate, 2% paraformaldehyde, 75 mM t-Lysin dihydrochloride, and 37.5 mM sodium phosphate, and then stained in a filtered solution of 0.3% oil red O in 60% isopropanol for 15 minutes. The red-stained, lipid vacuole–containing cells in a well were counted. To rescue osteoblast and adipocyte differentiation of PPARγ–/– ES cells, the recombinant retrovirus vector carrying the PPARγ gene and empty vector were constructed as previously described (22). ES cells were infected with equal titers of each recombinant virus as described (20), with some modification.

Skeletal morphology and blood chemistry. A bone radiograph was taken with a soft x-ray apparatus (SOFTEX, CMB-2, Tokyo, Japan). A three-dimensional CT scan was taken using a composite x-ray apparatus (SOFTEX; CMB-2, Tokyo, Japan). Incorporation of 1H]-thymidine (1 µCi/ml in the medium) was measured using a Wako ALP kit (Wako Pure Chemical Industry Ltd.) and the protein content was determined using a BCA protein assay reagent (Pierce Chemical Co., Rockford, Illinois, USA). The parameters for the trabecular bone culture, cells were sonicated in 10 mM Tris-HCl buffer (pH 8.0) containing 1 mM MgCl2 and 0.5% Triton X-100. ALP activity in the lysate was measured using a Wako ALP kit (Wako Pure Chemicals Industry Ltd.), and the protein content was determined using a BCA protein assay reagent (Pierce Chemical Co., Rockford, Illinois, USA). The primary bone marrow cell cultures. Bone marrow cells were collected from long bones of 8-week-old PPARγ–/– and WT male littermates. Cells were plated at a density of 106 cells on a six-multiwell plate in α-MEM containing 10% FBS, with 1 µM troglitazone for the adipogenesis assay and with 50 µg/ml ascorbic acid and 10 mM β-glycerophosphate for osteogenesis assay. The oil red O staining was performed as mentioned above at 14 days of culture. For the alkaline phosphatase (ALP), cultured plates were rinsed with PBS, fixed in 100% ethanol at 10 days of culture, and stained with Tris-HCl–buffered solution (pH 9.0) containing naphtholph AS-MX phosphate as a substrate and Fast Blue BB salt (Sigma-Aldrich, St. Louis, Missouri, USA) as a coupler. For the Alizarin red S staining, cultured plates were rinsed with PBS at 21 days of culture, fixed in 10% buffered formalin, and stained with 2% Alizarin red S (pH 4.0) (Sigma-Aldrich). The von Kossa staining was performed as mentioned above at 28 days of culture. For the growth curve assay, bone marrow cells derived from either WT or PPARγ–/– littermates were inoculated at a density of 105 cells per dish in 10-cm culture dishes in α-MEM containing 10% FBS, 50 µg/ml ascorbic acid, and 10 mM β-glycerophosphate, and then cultured for 3 days. The adherent cells were then harvested and inoculated at a density of 3 × 104 cells/dish in 10-cm culture dishes and further cultured in the same medium. The number of cells per dish was counted 1, 2, 3, and 4 days after the seeding.

Real-time quantitative RT-PCR. Total RNA was extracted with an ISOGEN kit (Wako Pure Chemicals Industry Ltd., Osaka, Japan), according to the manufacturer’s instructions, from ES cells cultured for 10 days after the embryoid bodies were transferred to gelatinized plates and from bone marrow cells cultured for 14 days after the seeding. One microgram of RNA was reverse-transcribed using a Takara RNA PCR Kit (AMV) version 2.1 (Takara Shuzo Co., Shiga, Japan) to make single-stranded cDNA. PCR was performed on an ABI Prism 7000 Sequence Detection System (Applied Biosystems Inc., Foster City, California, USA). The PCR reactions consisted of Quantitect SYBR Green PCR Master Mix (QIAGEN, Tokyo, Japan), 0.3 µM specific primers, and 500 ng of cDNA. Relative levels of mRNA of a specific gene were calculated using the standard curve generated with cDNA dilutions, with normalization to actin as an internal control. PCR primers of specific genes used for amplification are available upon request.

Primary osteoblastic cell cultures. Osteoblastic cells were isolated from calvariae of neonatal WT and PPARγ–/– littermates as previously described (27). For the cell proliferation assay, cells were inoculated at a density of 104 cells/well in a 24-multiwell plate, cultured in the same medium for 48 hours, and deprived of serum for 12 hours before adding the experimental medium with and without troglitazone (1 µM) or FGF-2 (1 nM; Kaken Pharmaceutical Co., Chiba, Japan). Incorporation of [3H]-thymidine (1 µCi/ml in the medium) added for the final 3 hours was measured after 24 hours of culture. For ALP activity measurement, cells were inoculated at a density of 104 cells/well in a 24-multiwell plate and cultured in α-MEM containing 10% FBS and 50 µg/ml ascorbic acid with and without troglitazone (1 µM) or bone morphogenetic protein-2 (BMP-2; 30 ng/ml; Yamanouchi Pharmaceutical Co., Tokyo, Japan). At 14 days of culture, cells were sonicated in 10 mM Tris-HCl buffer (pH 8.0) containing 1 mM MgCl2, and 0.5% Triton X-100. ALP activity in the lysate was measured using a Wako ALP kit (Wako Pure Chemicals Industry Ltd.), and the protein content was determined using a BCA protein assay reagent (Pierce Chemical Co., Rockford, Illinois, USA). For Alizarin red S and von Kossa stainings, cells were inoculated at a density of 5 × 104 cells/well in a six-multiwell plate in α-MEM containing 10% FBS, 50 µg/ml ascorbic acid, and 10 mM β-glycerophosphate.
erophosphate, and were stained at day 21 and 28, respectively, as mentioned above. Difference in maturity between the bone marrow cells and the calvarial osteoblasts was examined by the calcified nodule formation determined by Alizarin red S staining and the osteocalcin expression determined by real-time PCR analysis.

Assays for osteoclastic cells. Tartrate-resistant acid phosphatase-positive (TRAP-positive) multinucleated osteoclasts were generated by coculturing osteoblastic cells (10^4 cells/well) and bone marrow cells (5 × 10^5 cells/well) derived from either WT or PPAR γ+/– litters, as mentioned above, in a 24-multiwell plate in α-MEM containing 10% FBS for 6 days with and without 1,25(OH)2D3 (10 nM), prostaglandin E2 (100 nM), and IL-11 (10 ng/ml). Cells positively stained for TRAP containing more than three nuclei were counted as osteoclasts. To determine bone resorption activity, osteoclasts formed by the coculture on 0.24% collagen gel coated on 100-mm dishes were digested with 0.2% collagenase solution, and a 1:50 aliquot including osteoclasts was seeded on a dentine slice. After 48 hours of culture in α-MEM containing 10% FBS, the total area of pits stained with 0.5% toluidine blue was evaluated using an image analyzer. To study the role of PPAR γ intrinsic to osteoclastic cells, we used the M-CSF–dependent bone marrow macrophage (M-BMMφ) culture system as described previously (28). Briefly, bone marrow cells from WT or PPAR γ−/− mice were seeded at a density of 3 × 10^5 cells/well in a 24-multiwell plate and cultured in α-MEM containing 10% FBS with M-CSF (100 ng/ml). After culturing for 3 days, adherent cells (M-BMMφ) were further cultured with M-CSF (100 ng/ml) and soluble receptor activator of nuclear factor κB ligand (RANKL) (100 ng/ml) for 3 days. TRAP-positive osteoclasts were counted. To determine the survival, osteoclasts generated as above were deprived of M-CSF/soluble RANKL and cultured for an additional 48 hours. At 3, 6, 12, 24, and 48 hours, the TRAP-positive and trypan blue–negative osteoclasts were counted.

Statistical analysis. Means of groups were compared by ANOVA, and significance of differences was determined by post-hoc testing using Bonferroni’s method.

Results

Osteogenesis is enhanced in the homozygous PPAR γ-deficient ES cell culture. To examine the involvement of the PPAR γ signaling in fat and bone metabolism, we first compared the adipogenesis and osteogenesis in

Figure 1

Adipogenesis and osteogenesis in the mouse ES cell cultures of homozygous PPAR γ-deficient (PPAR γ–/–) and WT genotypes. As a rescue experiment, PPAR γ was reintroduced into PPAR γ–/– ES cells using a retrovirus vector carrying the PPAR γ gene (Rx-PPAR γ) or the same retrovirus vector without the PPAR γ gene (Rx-vector) as a control. (A) The upper row shows the adipogenesis determined by the oil red O staining of the ES cell culture in DMEM/10% FBS with troglitazone. The number of oil red O–positive cells stained in red was counted and shown in the left graph as the cells per square centimeter. The images in the lower row indicate the osteogenesis determined by the von Kossa staining of the ES cell culture in DMEM/10% FBS without any osteogenic supplements. The number of von Kossa–positive calcified nodules stained in black was counted and shown in the right graph as the number per square centimeter. Scale bar: 20 μm. (B) Relative mRNA levels of the marker genes for osteoblasts — COL1A1, osteocalcin and Runx2 — determined by real-time quantitative RT-PCR 10 days after the embryoid bodies were transferred to a gelatinized six-multiwell plate in DMEM/10% FBS without any osteogenic supplements. The ordinate axis indicates the relative amount of mRNA as compared with that of WT. Data are expressed as means (bars) ± SEMs (error bars) for eight wells per group. *Significant difference from the WT culture, P < 0.01. †Significant restoration by Rx-PPAR γ as compared with the control PPAR γ–/– and PPAR γ–/– plus Rx-vector cultures; P < 0.01. Cont, control.
the cultures of ES cells between PPARγ–/– and WT (WT or PPARγ+/−) genotypes isolated from blastocysts generated by mating PPARγ+/− mice (Figure 1A). In the presence of troglitazone, a thiazolidinedione that is a potent ligand of PPARγ, a substantial amount of oil red O–positive adipocytes was formed from WT ES cells, whereas adipogenesis was not seen in the PPARγ–/– ES cell culture (Figure 1A, upper row of photographs). To confirm the direct association between PPARγ and adipogenesis, PPARγ was reintroduced into PPARγ–/– ES cells using a retrovirus vector carrying the PPARγ gene (Rx-PPARγ). Adipogenesis was restored to the level similar to that of WT culture, although introduction of the same retrovirus vector without the PPARγ gene (Rx-vector) altered neither (Figure 1A, lower row of photographs). Quantitative analysis of the mRNA levels by the real-time RT-PCR method revealed that the marker genes for osteoblasts — type I collagen α1 chain (COL1A1), osteocalcin, and Runx2 — were upregulated in the PPARγ–/– ES cell culture as compared with the WT culture (Figure 1B). Reintroduction of PPARγ into the PPARγ–/– culture by Rx-PPARγ significantly decreased the nodule formation and osteogenic marker gene expressions, while the control Rx-vector altered neither (Figure 1, A and B). When one takes these results together, the observed mirror image regulations between adipogenesis and osteogenesis by loss and gain of the PPARγ function suggest a switching mechanism between the two differentiation pathways from common progenitors through the PPARγ signaling.

PPARγ haploinsufficiency leads to high bone mass in vivo. To learn the effect of the PPARγ insufficiency in vivo, we analyzed the bones of PPARγ+/– mice because the homozygous deficient fetuses died too early for their skeletal analyses to be performed. PPARγ+/− mice showed normal weight gain without visible general lipodystrophy on a standard diet during the observation period of up to 52 weeks of age. The lengths of the trunk and long bones were also similar to those of WT littermates, indicating that PPARγ is not involved in the regulation of skeletal growth. X-ray and three-dimensional CT analyses of femora and tibiae, however, revealed that PPARγ–/– mice showed about 40% higher trabecular bone mass than WT littermates at 8 weeks of age (Figure 2, A and B).
Contrarily, the number of adipocytes in the bone marrow determined as described below tended to be lower in the \( PPAR_\gamma^{+/-} \) long bones compared with WT (Figure 2B). Similar changes of bone and fat were also seen in vertebral bodies (data not shown). To examine the involvement of systemic factors that are known to be related to bone and fat metabolism, the serum levels of insulin and leptin were compared between the two mouse genotypes. \( PPAR_\gamma^{+/-} \) mice showed lower, although not significantly lower, serum insulin level and higher leptin level than WT littermates as reported previously (22, 29). Since insulin is known to be osteogenic (30), whereas leptin is antiosteogenic (11, 13, 31), neither of the changes in these hormones could explain the increased bone mass in \( PPAR_\gamma^{+/-} \) mice.

Because age-related osteoporosis is known to be accompanied by reciprocal increase of bone marrow adipocytes (7–9), we further compared the bones of \( PPAR_\gamma^{+/-} \) and WT littermates at 52 weeks of age (Figure 2, C and D). The BV of femora and tibiae was decreased in both mouse genotypes at this age as compared with that at 8 weeks; however, the difference of BV between \( PPAR_\gamma^{+/-} \) and WT became more prominent at 52 weeks than at 8 weeks (95% versus 40%, respectively). The number of bone marrow adipocytes, which are shown as oval vacuoles by the toluidine blue staining (Figure 3A, right), was significantly decreased in \( PPAR_\gamma^{+/-} \) mice at this age. This tendency was similarly observed in vertebral bodies (data not shown). Both insulin and leptin levels at this old age showed patterns similar to those at 8 weeks, although significant differences between the genotypes were not seen.

\( PPAR_\gamma \) haploinsufficiency leads to osteoblastogenesis in vivo. We further performed histological analyses of the proximal tibiae of 8-week-old \( PPAR_\gamma^{+/-} \) mice. Villanueva-Goldner staining indicated increases of trabecular bones stained in green and osteoid surface stained in red in \( PPAR_\gamma^{+/-} \) mice as compared with WT littermates; however, bone formation by individual osteoblasts determined by the calcein double labeling and the number of TRAP-positive osteoclasts was similar for the two groups (Figure 3A). Bone histomorphometric analyses (Figure 3B) confirmed the increase of BV by \( PPAR_\gamma \) haploinsufficiency to be about 40%. Among bone formation parameters, osteoblast surface and osteoid surface, both representative of the number of osteoblasts, were more than double in \( PPAR_\gamma^{+/-} \) than in WT littermates, while the mineral apposition rate that reflects the bone formation ability of individual osteoblasts did not differ between them. Consequently, bone formation rate that is determined by the number and the function of osteoblasts became about twice that by \( PPAR_\gamma \) haploinsufficiency. Bone resorption parameters, osteoclast number, and eroded surface did not differ between \( PPAR_\gamma^{+/-} \) and WT mice. Taking these histologi-
cal observations together, *PPARγ*+/– mice exhibited high bone mass with increased osteoblastogenesis but normal osteoblast and osteoclast functions. The thickness of the growth plate at the proximal tibiae was not different between *PPARγ*+/– and WT littermates (80.4 ± 9.6 and 82.7 ± 10.3 µm, mean ± SEM of eight mice each, respectively), confirming that *PPARγ* signaling is not important for bone growth through chondrocyte functions.

*PPARγ* haploinsufficiency does not affect bone loss by OVX. To investigate the involvement of the *PPARγ* signaling in the mechanism of bone loss by estrogen deficiency, OVX or sham operation was undertaken on 26-week-old female *PPARγ*+/– and WT littermates, and BV was compared 4 weeks after the operation. X-ray and three-dimensional CT analyses of femora and tibiae suggested that both *PPARγ*+/– and WT mice showed similar bone loss by OVX (Figure 4A). Histomorphometric analyses (Figure 4B) showed that BV was about 30% decreased by OVX in both *PPARγ*+/– and WT mice. These decreases were accompanied by increases in bone formation and bone resorption parameters, indicating a state of high bone turnover, in both genotypes. Hence, *PPARγ* haploinsufficiency did not affect the change of bone metabolism induced by OVX, suggesting that the *PPARγ* signaling does not contribute to osteopenia caused by estrogen deficiency.

*PPARγ* haploinsufficiency leads to osteoblastogenesis from cultured bone marrow cells. To investigate the cellular mechanism underlying the abnormality in the bone of *PPARγ*+/– mice, ex vivo cultures of bone marrow cells derived from long bones were performed. We first compared the cell proliferation determined by the growth curve for 4 days and found no difference between *PPARγ*+/– and WT marrow cultures (Figure 5A). Adipogenesis from marrow cells in the presence of troglitazone was confirmed to be inhibited by *PPARγ* haploinsufficiency, however, since the number of oil red O–positive adipocytes was decreased in the *PPARγ*+/– culture to about half that of the WT culture (Figure 5B). We further examined osteoblastogenesis in the bone marrow cell culture by comparing the numbers of colonies positively stained with ALP, Alizarin red S, and von Kossa (Figure 5C). All colonies were markedly increased in the *PPARγ*+/– culture as compared with the WT culture, indicating the increase of osteoblastogenesis from bone marrow progenitors by *PPARγ* haploinsufficiency.

To further investigate the regulation of expression of genes related to bone metabolism by *PPARγ* haploinsufficiency, we compared mRNA levels of key or marker molecules for adipocyte and osteoblast differentiations between *PPARγ*+/– and WT bone marrow cells (Figure 5D). As expected, *PPARγ* expression was reduced in the *PPARγ*+/– marrow cells compared with the WT. The levels of expression of other key factors for adipocyte differentiation, C/EBP-β and C/EBP-δ, in the *PPARγ*+/– marrow cells were comparable to those of WT, indicating that *PPARγ* was not essential for induction of these C/EBPs; contrarily, C/EBP-α was significantly reduced. Based on previous observations (16, 17, 22, 32, 33) and these results, it appears that C/EBP-β and C/EBP-δ lie upstream of *PPARγ*, while C/EBP-α is...
regulated, at least in part, downstream of PPARγ. Regarding osteogenic factors, expressions of the putative central determinants of major pathways for osteoblast differentiation, Runx2 (34), osterix (35), and LDL receptor–related protein 5 (LRP5) (36), were increased in the PPARγ+/− culture as compared with WT, indicating that the PPARγ signaling directly or indirectly impacts these major pathways for osteoblast differentiation. Expressions of matrix proteins representing osteogenesis, COL1A1, ALP, osteocalcin, and osteopontin, were also higher in the PPARγ+/− culture than in the WT culture, which was consistent with the in vivo histomorphometric data showing high bone mass with increased osteoblastogenesis.

When we examined the proliferation, differentiation, and matrix synthesis of cultured calvarial osteoblasts, which we confirmed to be more mature than bone marrow cells, none of them showed a difference between PPARγ+/− and WT mice (data not shown). This indicates that PPARγ haploinsufficiency affects only marrow progenitors, but not cells that are more committed to osteoblastic lineage. Furthermore, studies using the coculture system of marrow cells/calvarial osteoblasts and the M-BMMφ culture system (28, 37) also failed to show difference of differentiation, bone-resorbing activity, or survival of the osteoclastlike cells, suggesting that PPARγ is not important for osteoclast functions.

Discussion
Osteoblastogenesis was upregulated not only in the PPARγ−/− bone in vivo, but also in the cultures of PPARγ−/− ES cells and PPARγ+/−.
primary bone marrow cells. Considering that the former culture was performed in the absence of any osteogenic stimulation, under which condition no WT stem cells can differentiate into osteoblasts, the intrinsic PPARγ signaling seems to function as a potent suppressor of commitment and differentiation to the osteoblastic lineage. Its molecular mechanism remains unclear, however. A previous report showed that a stable transfection of PPARγ and its activation with a thiazolidinedione-suppressed Runx2, type I collagen, and osteocalcin syntheses in the culture of a stromal cell line (38). Although the present study also showed that steady-state mRNA levels of the key molecules for osteoblast differentiation, Runx2, osterix, and LRP5, were upregulated in primary cultured marrow cells with PPARγ haploinsufficiency (Figure 5D), whether this is transcriptional regulation or secondary to the increase in cells of osteoblast lineage in the culture is unknown. The predicted region of the PPARγ-responsive element PPRE (TGACCTnnTGACCT) has not been identified in the promoter of these genes and was not found by our genomic search, at least in the region between 4.0 kb upstream and 0.5 kb downstream of the transcriptional start point of runx2 (GenBank accession number NT 039655), osterix (NT 039621), and lrp5 (NT 039684) genes. It should, however, also be noted that several reports have indicated that PPARγ regulates gene expression independently of PPRE, that is, by interfering with the function of AP-1, signal transducer and activator of transcription 1 (STAT1), and NF-kB (39), or by inhibiting the function of GHP-1, a transcription factor implicated in pituitary-specific gene expression (40). The AP-1 family members may possibly play a role in the mechanism, especially for mesenchymal cells. Activation of PPARγ is reported to suppress c-fos expression (41). Another AP-1 family member, DeltaFosB, is known to be a positive regulator of osteoblast differentiation, and the transgenic mouse leads to postnatal high bone mass with increased osteoblastogenesis and decreased adipogenesis in bone marrow (12). Another possible molecular mechanism is the interaction of PPARγ with the TGF-β/Smad3 signaling, which inhibits osteoblast differentiation (42, 43). Since Smad3 is reported to interact physically with Runx2 (42) and PPARγ (44), the interference by PPARγ with the Smad3 inhibition of Runx2 might be involved in the switching mechanism between adipocyte and osteoblast differentiation. The interaction of PPARγ with Wnt signaling might be another issue to pursue. The canonical Wnt pathway, likely mediated by Wnt10b, is known to maintain preadipocytes in an undifferentiated state through stabilization of cytosolic β-catenin (45, 46). Since activation of PPARγ with troglitazone is not sufficient to repress expression of Wnt10b, Wnt signaling might lie upstream of PPARγ. Recently, in addition to LRPs, which is a coreceptor of Wnt, the canonical Wnt-signaling molecules β-catenin and glycogen synthase kinase-3β have been reported to stimulate osteoblast differentiation (47, 48), indicating a switching between adipogenesis and osteogenesis by the Wnt signaling. Further studies on functional interaction of PPARγ with the transcriptional and signaling molecules above will elucidate the switching mechanism between the two differentiation pathways from common progenitors.

PPARγ may also inhibit osteogenesis indirectly through its stimulation of adipogenesis from marrow progenitors that can give rise to either osteoblasts or adipocytes. In fact, many experimental models have provided substantial evidence for this reciprocal relationship between cell lineages (6, 10–15), and there is little doubt that adipogenesis increases as BV decreases, suggesting that marrow adipogenesis has important implications in osteogenic disorders (7–9). Evidence of the transdifferentiation of stromal cells actually suggests a large degree of plasticity between osteoblasts and adipocytes (4, 49), although it is not clear at what point the phenotype of these multipotent cells becomes committed to either osteoblast or adipocyte differentiation. Since differentiated osteoblasts indicated by osteocalcin expression are reported to undergo adipogenic differentiation (4), it is possible that the reciprocal relationship between osteogenesis and adipogenesis may, at least in part, be due to the transdifferentiation between rather differentiated cells of the two lineages. To determine the role of PPARγ in more differentiated osteoblastic cells than bone marrow cells, we used calvarial cells whose spontaneous differentiation is known to follow not only the osteogenic pathway but also the adipogenic pathway (49). Despite the existence of PPARγ expression in these cells as well, its haploinsufficiency did not affect the cell functions, suggesting that PPARγ signaling may be involved in the earlier, but not the later, stage of relationship between the two cell lineages.

Hormones regulating bone and fat metabolisms include insulin and leptin, both of which are known to be related to the PPARγ signaling. Insulin is known to play important anabolic roles in bone (30), and deficiency of insulin signaling is associated with osteopenia both in mice and humans (27, 50–52). A series of reports demonstrated that leptin, a well-known anorexigenic hormone secreted by adipocytes (53), also shows antiosteogenic action centrally through the hypothalamic and sympathetic nervous systems (11, 13, 31). In the present study, neither insulin nor leptin seemed to mediate the high bone mass in PPARγ−/− mice, since the serum levels of these hormones were, quite unexpectedly, the opposite of those causing osteogenic functions. PPARγ activation is known to cause insulin sensitivity, thus PPARγ−/− mice were assumed to develop insulin resistance; however, the serum insulin level was normal or somewhat decreased as previously reported (22, 29). This appears, at least in part, due to hypersecretion of leptin, which was also unexpected, given that the marrow adipocytes, a positive regulator of leptin expression, were decreased in PPARγ−/− mice. Our previous studies clearly demonstrated that cultured primary adipocytes from PPARγ−/− mice expressed and secreted increased levels of leptin as compared with those from WT (22, 29). In this respect, since leptin is known to have a functional PPRE whose activity is suppressed by PPARγ activation in adipocytes (54, 55), it is likely that the increased level of leptin is due to a partial release from the suppressive effect of PPARγ on leptin gene transcription by loss of one PPARγ allele.

Age-related bone loss has been suggested to be attributable to increased adipogenesis at the expense of osteoblastogenesis (7–9). Indeed, studies of SAMP6 mice, a murine model of age-related osteopenia, have established a tight association between osteopenia and enhanced adipogenesis (10, 15). The fact that the effects of PPARγ haploinsufficiency on both the increase in bone volume and the decrease in adipocytes were stronger at 52 weeks than at 8 weeks suggests the involvement of the PPARγ signaling in the pathophysiology of human age-related osteoporosis. In fact, our preliminary examination of the bone marrow specimen from patients with femoral neck fracture actually showed increases of both the PPARγ mRNA level and fat mass in older patients (data not shown) as compared with those in younger patients, although the causality between PPARγ level and adipogenesis remains unknown since adipocytes can be both the source and the target...
of PPARγ. An association study between bone density and a genetic polymorphism of PPARγ in postmenopausal women implies the involvement of PPARγ in bone loss, although the functional relevance remains unclear (56). We should, however, keep in mind that there are two distinct factors that determine involutional osteoporosis: a rapid bone loss after menopause as a result of estrogen withdrawal and a gradual age-related bone loss thereafter (57). From the present study showing that PPARγ insufficiency did not affect bone loss by OVX, PPARγ may not be involved in the former stage, but may play a role in the latter. To confirm the involvement of PPARγ in human osteoporosis, the next task ahead of us will be to perform a genetic association study with stratified analysis by age and menopausal state, and more importantly, to use a cohort population.

We conclude herein that PPARγ haploinsufficiency leads to the increase of bone mass by stimulating osteoblastogenesis from bone marrow progenitors without affecting differentiated osteoblasts or osteoclast lineage cells. Based on the present and previous evidence presented, we believe that PPARγ may be a novel target for therapeutic intervention of osteopenic disorders, although the mechanism remains to be clarified. Appropriate functional antagonism of PPARγ may provide a potentially novel approach to increasing bone formation and therefore, as a stand-alone therapy or in combination with an antiresorptive medication, may provide more efficacious prevention or treatment of osteoporosis.

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