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Genetic ablation of IL-6 in gp130Y757F/Y757F mice exacerbated this osteopenia by reducing ObS/BS without affecting OcS/BS. Thus, while IL-6 is critical for high bone formation in gp130Y757F/Y757F mice, it is not involved in the increased osteoclastogenesis. In conclusion, gp130 is essential for normal bone growth and trabecular bone mass, with balanced regulation depending on selective activation of STAT1/3 and SHP2/ras/MAPK, respectively. Furthermore, the latter pathway can directly inhibit osteoclastogenesis in vivo.

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Glycoprotein 130 regulates bone turnover and bone size by distinct downstream signaling pathways

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The gp130-dependent cytokines, which signal through at least two intracellular pathways, regulate osteoclast and osteoblast formation. To define their roles in regulating bone mass, we analyzed mice in which gp130 signaling via either the signal transducer and activator of transcription (STAT) 1/3 (gp130∆STAT/∆STAT) or SHP2/ras/MAPK (gp130∆STAT/∆STAT) pathway was attenuated. In gp130∆STAT/∆STAT mice, trabecular bone volume (BV/TV) and turnover were normal, but bone length was reduced by premature growth plate closure, indicating an essential role for gp130-STAT1/3 signaling in chondrocyte differentiation. In contrast, while bone size was normal in gp130∆STAT/∆STAT mice, BV/TV was reduced due to high bone turnover, indicated by high osteoclast surface/bone surface (OcS/BS) and osteoblast surface/bone surface (ObS/BS). Furthermore, generation of functional osteoclasts from bone marrow of gp130∆STAT/∆STAT mice was elevated, revealing that while gp130 family cytokines stimulate osteoclastogenesis through the osteoblast lineage, gp130, via SHP2/Ras/MAPK, inhibits osteoclastogenesis in a cell lineage–autonomous manner. Genetic ablation of IL-6 in gp130∆STAT/∆STAT mice exacerbated this osteopenia by reducing ObS/BS without affecting OcS/BS. Thus, while IL-6 is critical for high bone formation in gp130∆STAT/∆STAT mice, it is not involved in the increased osteoclastogenesis. In conclusion, gp130 is essential for normal bone growth and trabecular bone mass, with balanced regulation depending on selective activation of STAT1/3 and SHP2/ras/MAPK, respectively. Furthermore, the latter pathway can directly inhibit osteoclastogenesis in vivo.


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Nonstandard abbreviations used: leukemia inhibitory factor (LIF); cardiotoxin (CT-1); ciliary neurotrophic factor (CNTF); LIF receptor (LIF-R); OsM receptor (OsM-R); signal transducer and activator of transcription (STAT); proliferative zone width (Prol.Z.Wi); hypertrophic zone width (Hyp.Z.Wi); microtome (micro-CT); parathyroid hormone (PTH); bone marrow macrophage precursor (BMMP); receptor activator of NF-κB ligand (RANKL); trartrate-resistant acid phosphatase (TRAP); multinucleated cell (MNC); MEM; α modification (α-MEM); growth plate width (G.P.Wi); growth plate closure (G.P.Cl); trabecular bone volume (BV/TV); trabecular number (Tb.N); trabecular thickness (Tb.Th); osteoclast surface/bone surface (OcS/BS); osteoblast surface/bone surface (ObS/BS); osteoid volume/bone volume (OV/BV); mineral appositional rate (MAR).

Introduction

The gp130 family of cytokines includes leukemia inhibitory factor (LIF), IL-11, IL-6, cardiotrophin-1 (CT-1), oncostatin M (OsM), and ciliary neurotrophic factor (CNTF). These cytokines signal by forming a receptor complex containing the common gp130 coreceptor signaling subunit (1). The specific components of this receptor complex depend on the nature of the bound ligand. IL-11 and IL-6 bind to specific α-receptor subunits that lack their own signaling domains and are entirely dependent for intracellular signaling on the function of gp130 through gp130 β-subunit homodimers (1). In contrast, LIF, OsM, CT-1, and CNTF signal through gp130 heterodimers containing either LIF receptor (LIF-R) or OsM receptor (OsM-R) β subunits. Formation of either gp130-containing complex results in activation of cytoplasmic Janus kinases, which phosphorylate tyrosine residues on gp130, LIF-R, and OsM-R β (2). Dimerization of gp130 results in activation of the signal transducer and activator of transcription (STAT) 1 and STAT3 (3, 4) and the SHP2/ras/MAPK (5) signaling pathways. The use of distinct downstream signaling pathways from different intracellular regions of the same gp130 receptor subunit enables ligand- and tissue-specific activation of distinct
sets of target genes and, thus, distinct physiologic effects of the gp130 cytokines in vivo (6).

The coordinated activities of osteoclasts (bone-resorbing cells) and osteoblasts (bone-forming cells) determine the overall structure and strength of the mammalian skeleton. LIF, IL-11, IL-6, CT-1, and OsM have all been reported to stimulate osteoclastogenesis (7–9) and, in some cases, to stimulate osteoblast proliferation or differentiation (10–12) in cell culture systems. Transgenic overexpression of IL-11 results in increased osteoblast generation in vivo and ex vivo (13), yet overexpression of IL-6 leads to low levels of both osteoblast and osteoclast formation (13, 14). Furthermore, studies of knockout mice revealed elevated osteoclast numbers in the bones of LIF-Rβ knockout mice (15), but low bone resorption and formation in the absence of IL-11R (16). Despite the shared use of receptor subunits, these cytokines are therefore likely to mediate their effects on different bone cell types through different intracellular signaling pathways.

Both bone formation and resorption are altered significantly in gp130−/− mice (17, 18). Although this mutation results in neonatal lethality, reduced trabecular bone mass is a feature of the neonatal skeleton of these mice. The finding of increased osteoclastogenesis both in vivo and in ex vivo cultures was surprising, given the previously reported capacity of gp130 cytokines to stimulate osteoclast formation in vitro (7–9). Furthermore, in wild-type cultures, osteoclastogenesis was inhibited by a gp130-neutralizing antibody in vitro (9), and bone loss in ovariectomized mice was prevented by treatment of the mice with a gp130 nonpeptide antagonist (19). The gp130−/− mice also demonstrated shortened limbs (17, 18); these bones are formed by controlled growth and mineralization of a cartilage model (endochondral ossification), and this defect implies an impairment in chondrocyte differentiation in the absence of gp130.

In this study, we sought to determine the specific contributions made by the two signaling pathways emanating from gp130 to normal osteoclastogenesis and to endochondral ossification by utilizing two gp130 “knock-in” mouse strains homozygous for mutations in gp130 at either the SHP2/ras/MAPK or the STAT1/3 activation sites. Specifically, gp130ΔSTAT/ΔSTAT mice express a knock-in mutant of gp130 with a C-terminally deleted gp130 that lacks all STAT1/3 binding and activation sites while retaining the capacity to activate SHP2/ras/MAPK (20). Conversely, gp130Y757F/Y757F mice express gp130 with a point mutation that selectively blocks the SHP2/ras/MAPK pathway, while STAT1/3 signaling remains intact (6).

Methods

Animals. Animals were generated as described previously (6, 20). Briefly, gp130ΔSTAT/ΔSTAT mice express a knock-in mutant of gp130 with a C-terminal deletion of the STAT1/3 binding and activation sites (20), while gp130Y757F/Y757F mice express gp130 with a point mutation that selectively blocks the SHP2/ras/MAPK pathway (6). Compound gp130Y757F/Y757F:IL-6−/− mice were generated from gp130Y757F/Y757F mice and IL-6−/− mice (21). All mouse strains were kept on a mixed 129/B6 background. For each experiment, littermate controls were used. All animal handling procedures were approved by the animal ethics committees at the Ludwig Institute for Cancer Research (Parkville, Victoria, Australia) and/or St. Vincent’s Health (Fitzroy, Victoria, Australia).

Histomorphometry. Tibiae were collected from 4-week-old, 12- to 16-week-old, or 6-month old mice, fixed in 4% paraformaldehyde in PBS, and embedded in methylmethacrylate (22). Double-fluorochrome labeling was performed with calcine injections 10 and 3 days before tissue collection, as described previously (22). Sections 5-μm-thick sections of the proximal tibia were stained with toluidine blue or analyzed unstained for fluorochrome labels according to standard procedures using the Osteomeasure system (OsteoMetrics Inc., Decatur, Georgia, USA). Tibial cortical thickness and periosteal mineral appositional rates were measured as described previously (22). Femoral length and width were determined from contact x-rays that were scanned and measured using NIH Image 1.62 as described previously (22). Growth plate width and thickness of the hypertrophic and proliferative zones were measured as described previously using the Osteomeasure system (OsteoMetrics Inc.) (22); proliferative zone width (Prol.Z.Wi) was defined as the width of the growth plate area in which proliferating chondrocytes were observed, and the hypertrophic zone width (Hyp.Z.Wi) was defined as the width of the growth plate in which whole lacunae containing hypertrophic chondrocytes were observed. In the gp130ΔSTAT/ΔSTAT mice, where regions of the growth plate were completely closed, these parameters were measured only in those regions where cartilage was present. In addition, length of growth plate closure (complete absence of cartilage) was measured as a percentage of total growth plate length across the tibia for all genotypes using the Osteomeasure system (OsteoMetrics Inc.).

Microtomography (micro-CT). Proximal tibial bone was assessed by micro-CT measurement (microCT-20; Scanco, Bassersdorf, Switzerland) on samples from male mice 8–12 weeks of age (n = 7 per group), starting 0.72 mm below the growth plate. A total length of 1.5 mm was measured at high resolution, yielding 168 sections 9 μm in thickness and a voxel resolution of 9 × 9 × 9 μm3. Threshold optimization was as detailed previously (23).

Serum biochemistry. Serum was collected from 10-week-old mice. Serum calcium was measured by reaction with o-cresolphthalein (Sigma-Aldrich, St. Louis, Missouri, USA). Intact parathyroid hormone (PTH) and serum IL-6 were measured by mouse-specific ELISAs (Immunotopics, San Clemente, California, USA, and Chemicon, Temecula, California, USA).

Cell cultures. The osteoclastogenic potential of bone marrow and bone marrow macrophage precursors
(BMMPs) was determined as described previously (24) by stimulation of bone marrow cell preparations flushed from the femora and tibiae of male and female mice (plated at a density of $10^5$ cells per 10-mm-diameter well) with soluble recombinant GST–receptor activator of NF-$\kappa$B ligand (RANKL) protein (25) and M-CSF (R&D Systems, Minneapolis, Minnesota, USA). Tartrate-resistant acid phosphatase (TRAP)-positive multinucleated cells (MNCs) were counted at day 7.

For resorption assays, bone marrow cells from $gp130^{\Delta STAT/\Delta STAT}$ and wild-type littermates were added to culture wells 6 mm in diameter ($10^5$ cells/well) containing 12-mm$^2$ dentine slices (26) and were stimulated with RANKL and M-CSF. After 14 days of incubation, the dentine slices were stripped of cells using 0.25 M NH$_4$OH, rinsed in water and ethanol, and then dried. For identification of resorption pits, the dentine slice surfaces were stained with xylene-free black ink and residual ink was removed by wiping the surface against absorbent paper. The proportion of the dentine surface resorbed was measured under transmission light microscopy by point counting (26).

Osteoblast differentiation was assessed with ex vivo osteoblast cultures derived from bone marrow precursors as described previously (27). Briefly, tibiae and femora from 8- to 10-week-old mice of each genotype were dissected free of adhering muscle. The ends were removed and the marrow cavity was flushed with MEM, $\alpha$-modification ($\alpha$-MEM). Cells were plated at a density of $2 \times 10^6$ cells/well in 24-well plates in $\alpha$-MEM containing 10% FCS, 10 nM dexamethasone, 10 mM $\beta$-glycerophosphate, and 50 mg/ml ascorbic acid. Cells were fixed on days 14, 21, and 28 and were stained for alkaline phosphatase to detect osteoblast colony formation or by a von Kossa technique to detect mineralization (28).

**Statistical analyses.** All data are presented as means ± SEM. Significant differences were determined by one-way or two-way ANOVA followed (where significant) by Fisher’s protected least significant difference post hoc.
test to identify the significant pairwise differences. Where data were not normally distributed (for percentage growth plate closure measurements), significant differences were identified by the nonparametric Kruskal-Wallis test. Simple linear regressions with ANOVA were used to determine the whether growth plate closure was associated with histomorphometric parameters reflecting growth plate chondrocyte differentiation of male and female gp130∆STAT/∆STAT mice. For all analyses, \( P < 0.05 \) was considered significant. All statistical analyses were carried out using StatView 5.0.1 (SAS Institute, Cary, North Carolina, USA).

**Results**

**Mild dwarfism due to premature growth plate closure in gp130∆STAT/∆STAT mice.** The most obvious phenotype detected in the gp130 mutants by x-ray analysis was a reduction in bone size in the gp130∆STAT/∆STAT mice (Figure 1a). Quantification revealed a significant reduction in both femoral length (12%; Figure 1b) and femoral width (18%; not shown) in both female and male gp130∆STAT/∆STAT mice. In contrast, gp130Y757F/Y757F mutant bones were normal in size (Figure 1b).

The reduced bone size in gp130∆STAT/∆STAT mice was associated with premature closure of the growth plate, the region in which chondrocyte proliferation usually determines longitudinal bone growth. In 16-week-old mice, where the zone of chondrocyte prolif-

| Table 1 Serum biochemistry and cortical bone histomorphometry |
|---------------------|---------------------|---------------------|
|                     | Wild-type           | gp130∆STAT/∆STAT    | gp130Y757F/Y757F   |
| Serum PTH (pg/ml)   | 33.1 ± 10.6         | 35.9 ± 20.0         | 54.5 ± 13.5        |
| Serum calcium (mg/dl)| 7.35 ± 0.12        | 8.22 ± 0.36         | 7.96 ± 0.23        |
| Serum IL-6 (pg/ml)  | 5.56 ± 1.82         | 8.40 ± 2.91         | 6.41 ± 1.52        |
| Tibial cortical thickness (\( \mu m \)) | 113.7 ± 3.0 | 105.8 ± 3.0 | 99.3 ± 3.1* |
| Periosteal MAR (\( \mu m \)/day) | 0.56 ± 0.10 | 0.73 ± 0.17 | 0.50 ± 0.13 |

Values are mean ± SEM from six to eight male mice per group at 14 weeks of age. *\( P < 0.001 \) vs. wild-type by one way ANOVA and Fisher’s post-hoc test.

**Figure 2**

Low trabecular bone volume in male and female gp130Y757F/Y757F mutant mice. (a) Representative images of von Kossa-stained tibiae of male 16-week-old WT, ∆/∆, and F/F mutant mice, showing differences in trabecular bone volume (white boxes show secondary spongiosa region used for histomorphometric measurements; note that this is a smaller anatomical region in the ∆/∆ mice due to the smaller bone size). Scale bar (blue), 500 \( \mu m \). Histomorphometric measures of BV/TV, Tb.N, and Tb.Th were all reduced in F/F mice compared with WT and ∆/∆ mice. (b) Micro-CT analysis and representative three-dimensional images of tibiae from male WT and F/F mice. (c) Representative images and histomorphometry for female 16-week-old WT and gp130 mutant mice. Scale bar (blue), 500 \( \mu m \). BV/TV is reduced in female F/F mice compared with WT and ∆/∆ mice. All values are mean ± SEM from a minimum of seven mice per group at 12–16 weeks of age for histomorphometry and 8–12 weeks of age for micro-CT. *\( P < 0.05 \); **\( P < 0.01 \); ***\( P < 0.001 \) vs. WT of the same sex.
High bone turnover and osteopenia in gp130Y757F/Y757F mice. Despite the reduced bone size, trabecular bone volume (BV/TV) in adult male and female gp130Y757F/Y757F mice was not significantly altered (Figure 2, a–c). In contrast, both male and female gp130Y757F/Y757F mice demonstrated a significant reduction in BV/TV, as initially suggested by two-dimensional histomorphometry and three-dimensional micro-CT analysis (Figure 2, a–c). In adult male gp130Y757F/Y757F mice, BV/TV and trabecular number (Tb.N) were halved, and trabecular thickness (Tb.Th) was significantly reduced, compared with wild-type littermates (Figure 2, a and b). In female mice of the same age, the effect was less dramatic and was not detected in Tb.Th, probably because, as is usual for female mice (29), BV/TV was already lower in females than in males (Figure 2c). Low BV/TV was also detected in 4-week-old and 6-month-old gp130Y757F/Y757F mice, while Hyp.Z.Wi did not. There was no change in either growth plate width or widths of either the hypertrophic or proliferating zones in gp130Y757F/Y757F mice (Figure 1 and results presented above). Neither cortical thickness nor periosteal apposition rate were altered in 16-week-old gp130Y757F/Y757F mice (Table 1), indicating that the small bone size in these mice did not relate to impaired periosteal growth. Our data therefore indicate that the presence of the gp130-dependent STAT1/3 signaling pathway, but not the SHP2/ras/MAPK pathway, plays an essential role in determining normal bone size by regulating chondrocyte proliferation.

Figure 3
Increased osteoclast formation in male and female gp130Y757F/Y757F mutant mice. Tibial trabecular OcS/BS was significantly increased in male (a) and female (b) F/F mice compared with WT and Δ/Δ mice. All values are mean ± SEM from a minimum of eight mice per group at 12–16 weeks of age. *P < 0.05; **P < 0.01; ***P < 0.001 vs. WT of the same sex. (c) Representative images of TRAP-stained primary spongiosa from WT and F/F mice. Note the large number of TRAP-positive osteoclasts (arrows) and the absence of trabecular bone in the F/F section. gp, growth plate; m, marrow; tb, trabecular bone. Scale bar (black), 200 µm. (d and e) In vitro osteoclastogenesis assays demonstrating increased TRAP-positive MNCs (TRAP+ MNCs/well) generated from RANKL-induced primary bone marrow cultures (d) and BMMP cultures (e) from WT and gp130 mutant mice. (f) Pit formation by osteoclasts generated from RANKL-induced primary bone marrow cultures from WT and gp130 mutant mice. All values are mean ± SEM of quadruplicate cultures from four separate mouse bone marrows of mixed sexes. *P < 0.05; **P < 0.01; ***P < 0.005 vs. WT culture.
compared with wild-type littermates (data not shown). These observations indicate that the gp130-dependent SHP2/ras/MAPK pathway is essential for the maintenance of normal trabecular bone structure.

The low BV/TV observed in the gp130Y757F/Y757F mice was associated with high osteoclast numbers in vivo. Within the secondary spongiosa, where bone is remodeled by repeated cycles of bone resorption and formation, the percentage of trabecular bone covered by osteoclasts (osteoclast surface/bone surface; OcS/BS) (Figure 3a) and the number of osteoclasts (not shown) were approximately doubled in the secondary spongiosa of male mice and, even in females, in which bone turnover is usually higher than in males (29), OcS/BS and osteoclast numbers were also increased when the SHP2/Ras/MAPK pathway was mutated (Figure 3b). In the gp130Y757F/Y757F mice, a large number of active osteoclasts was also detected at the base of the growth plate, where new trabecular bone is normally generated (Figure 3c). No change in osteoclast surface or number was detected in either male or female gp130STAT/STAT mice (Figure 3, a and b). Cortical thickness was also reduced in gp130Y757F/Y757F mice (Table 1).

Since the level of periosteal mineral apposition (transverse bone growth) was not altered in these mice (Table 1), this supports the finding of elevated osteoclast activity in gp130Y757F/Y757F mice.

Because elevated bone resorption in vivo can be associated with alterations in calcium homeostasis, we measured serum calcium and circulating PTH levels. Despite the high level of bone turnover, there was no significant alteration in either of these parameters in the gp130 mutants (Table 1), as has been observed in other animal models of high bone turnover osteopenia (30, 31), suggesting that the increased osteoclastogenesis is not secondary to altered PTH signaling or impaired calcium homeostasis.

Increased osteoclastogenesis in gp130Y757F/Y757F mice was reproduced in ex vivo cultures of bone marrow cells derived from these mice. While RANKL induced a similar level of osteoclast formation by wild-type and gp130STAT/STAT bone marrow, there was a significant elevation in osteoclastogenesis from RANKL-stimulated gp130Y757F/Y757F bone marrow (Figure 3d). When a more homogeneous preparation of osteoclast precursors from the marrow (BMMPs) was used, there was again a significant elevation in osteoclastogenesis from gp130Y757F/Y757F compared with wild-
The ability of these osteoclasts to resorb bone was confirmed by pit assays, demonstrating an increase in area resorbed by osteoclasts derived from gp130Y757F/Y757F marrow of a similar proportion to the increase in osteoclast numbers (Figure 3f). These data confirm that the defect in osteoclast number observed in vivo is not mediated by systemic hormones or altered paracrine signaling from osteoblasts but instead is autonomous within the hemopoietic cell lineage.

As observed in many osteopenic states, the high level of bone resorption in gp130Y757F/Y757F mice was associated with a high level of bone formation. Histomorphometric markers of bone formation were significantly elevated in both male and female gp130Y757F/Y757F mice. These included osteoblast surface/bone surface (ObS/BS), indicating a greater number of osteoblasts on trabecular bone surfaces; osteoid volume/bone volume (OV/BV), indicating a high level of new bone matrix (osteoid) production; and mineral appositional rate (MAR), indicating a high rate of mineralization of the bone matrix (Figure 4, a and b). We also observed increases in osteoid surface, osteoid thickness, and the number of osteoblasts on trabecular surfaces in both male and female gp130Y757F/Y757F mice (data not shown).

However, when osteoblast precursors were grown ex vivo from wild-type and gp130Y757F/Y757F bone marrow cells, we did not observe any significant difference in the ability of these precursors to differentiate into alkaline phosphatase–positive cells (osteoblasts) or any change in their ability to form mineralized nodules (Figure 4c). This indicated that the high level of bone formation in gp130Y757F/Y757F mice was not cell autonomous but may result from altered paracrine or endocrine activation taking place in the mutant mice. No change in any marker of bone formation was observed in gp130ΔSTAT/ΔSTAT mutants (Figure 4).

Discussion

The gp130-family of cytokines, including IL-6, IL-11, LIF, OSM, and CT-1, have been implicated in the determination of bone size and bone density from their effects in bone cell culture systems (9, 32, 33) and from the skeletal defects of transgenic (13, 14) and knockout mice (15, 16, 36). The contributions of the major intracellular gp130 signaling pathways responsible for bringing about these biological observations remain unknown. Here we have used knock-in gp130 mutant mice unable to elicit either gp130-dependent STAT1/3 or SHP2/ras/MAPK activation in order to detail the

Figure 5
IL-6 contributes to the high level of bone formation in gp130Y757F/Y757F mutant mice. (a) Representative images of von Kossa–stained tibiae from 16-week-old WT, F/F, and F/F:IL-6–/– (F/F:IL-6) mice. Scale bar (white), 500 µm. (b) BV/TV and Tb.N are reduced further in F/F:IL-6–/– mice compared with WT and F/F mice. Tb.Sp, trabecular separation. (c) Deletion of IL-6 rescues the high ObS/BS and reduces the OV/BV of F/F mutant mice but not the high Ocs/BS. All values are mean ± SEM from six female mice per group at 14–16 weeks of age. *P < 0.05; **P < 0.01; ***P < 0.001 vs. WT; **P < 0.01, *P < 0.05 vs. F/F.

Osteoclastogenesis phenotype is independent of IL-6, but that of the osteoblast is IL-6 dependent. Because IL-6 is known to stimulate osteoblast differentiation and osteoclastogenesis (32–35) and IL-6-stimulated STAT3 activation is enhanced in gp130Y757F/Y757F mice (6), we next set out to determine whether the increased bone formation and bone resorption in gp130Y757F/Y757F mice were mediated by IL-6. To do this, we carried out bone histomorphometry on adult gp130Y757F/Y757F:IL-6–/– mice. Surprisingly, the genetic ablation of IL-6, which itself does not lead to significant bone abnormality in the basal state (36), led to a further reduction in bone mass, indicated by BV/TV and Tb.N, as well as an increase in trabecular separation in compound mutant (gp130Y757F/Y757F:IL-6–/–) mice compared with single mutant gp130Y757F/Y757F:IL-6–/– mice, irrespective of sex (Figure 5 shows results from female mice only). While Ocs/BS remained high in the compound mutants, the high level of ObS/BS observed in gp130Y757F/Y757F:IL-6–/– mice was blocked and the high OV/BV was no longer significantly elevated in the absence of IL-6 (Figure 5), suggesting that IL-6 mediates the moderating effect of the gp130-dependent SHP2/ras/MAPK pathway on bone formation but not on osteoclastogenesis.
specific involvement of these pathways in regulating physiological bone turnover in vivo. This analysis allows us to define three distinct pathways that regulate bone cell function in vivo in response to gp130 family cytokines: (a) gp130-STAT1/3 regulation of chondrocyte proliferation, (b) gp130-SHP2/Ras/MAPK inhibition of osteoclastogenesis, and (c) IL-6 stimulation of osteoblasts through the gp130-STAT1/3 pathway (these pathways are summarized in Figure 6).

It has been suggested that gp130 family cytokines, particularly IL-6 and IL-11, may be involved in regulating sex steroid–mediated effects on bone growth and trabecular bone structure (12, 37). In this study, we observed an identical response in male and female mice to both gp130 mutations, suggesting that gp130 family members are not required for sex steroid–regulated differences in bone growth and bone turnover in the basal state.

Reduced bone size, premature growth plate closure, and reduced chondrocyte proliferation in gp130\(^{\text{ASTAT/ASTAT}}\) mice indicate that the gp130-dependent STAT1/3 signaling pathway plays an essential role in determining bone size, specifically by promoting growth plate chondrocyte proliferation. In gp130\(^{\text{ASTAT/ASTAT}}\) mice, regions of growth plate closure were detected as early as 12 weeks of age, which is very unusual, as it is rare to detect growth plate closure in mice even at 12 months of age.

It has been reported previously that the joints of some gp130\(^{\text{ASTAT/ASTAT}}\) mice exhibit excessive articular cartilage growth and joint stiffness (20); however, samples used in our study did not demonstrate this phenotype. Even if an early, undetected form of the articular cartilage defect were present in these mice, it is unlikely that this would also be a mechanism for growth plate closure (the opposite effect) in an area removed from the synovial environment. Indeed, in the only reported case we have found in which changes in cytokine levels at the joint have affected both articular and growth plate cartilage (38), the changes at the growth plate were restricted to the periosteal edge (the most external part of the growth plate). In contrast, regions of growth plate closure in the gp130\(^{\text{ASTAT/ASTAT}}\) mice occurred apparently at random across the width of the growth plate, with no growth plate closure being observed at the periosteal edge.

The mechanistic details of growth plate closure are not well understood in mammals, with part of the difficulty being that full growth plate closure, which is observed in adult humans, is not reported in rodents; thus, there is no good animal model available for the study of growth plate closure. Generally, it has been understood that growth plate closure occurs along with a gradual reduction in chondrocyte proliferation and is characterized by osteoclastic resorption of quiescent cartilage (39). Although reduced chondrocyte proliferation was observed in gp130\(^{\text{ASTAT/ASTAT}}\) mice and was correlated significantly with the extent of growth plate closure, we were surprised to find active chondrocyte proliferation immediately adjacent to regions of full growth plate closure (calcified bridge formation) in the gp130\(^{\text{ASTAT/ASTAT}}\) mice. This is consistent with a recent report of calcified bridge formation across the growth plate in rats as young as 3 months of age, although that study did not report the level of chondrocyte proliferation (40). Clearly, there is some mechanism whereby growth plate closure can occur without full cessation of chondrocyte proliferation, although we cannot say whether the effect observed in our study is physiological or pathological. Nevertheless, this suggests a heterogeneous response of the chondrocyte population to the absence of STAT1/3 signaling downstream of gp130.

**Figure 6**

Summary of the effects of gp130 signaling pathway mutations on bone structure and cell function. (a) WT mice, IL-6 stimulates osteoblast proliferation via the gp130-STAT1/3 pathway, and the osteoblasts in turn stimulate osteoclastogenesis by the RANKL pathway. Osteoclastogenesis is also inhibited by gp130 family cytokines, acting through the gp130-SHP2/Ras/MAPK signaling pathway within the hematopoietic lineage. Chondrocyte proliferation is stimulated by gp130 family cytokines through the gp130-STAT1/3 pathway. (b) In the absence of the gp130-induced STAT1/3 signaling pathway, indicated by the red X (in Δ/Δ mice), chondrocyte proliferation is no longer stimulated through the gp130-STAT1/3 pathway, leading to reduced bone length. Inhibition of osteoclastogenesis through the gp130-SHP2/Ras/MAPK pathway remains intact, and while IL-6 stimulation of osteoblast differentiation can no longer occur through the gp130-STAT1/3 pathway, this is clearly a minor pathway not required for the normal level of bone formation, as bone turnover and bone structure remain at the same level as in WT mice. (c) In F/F mice, IL-6-stimulated osteoblast proliferation via gp130-STAT1/3 is enhanced, leading to a greater number of osteoblasts. Deletion of the gp130-SHP2/Ras/MAPK pathway, indicated by the red X, releases inhibition, resulting in increased osteoclastogenesis. The increase in resorption appears to be greater than the increase in bone formation, leading overall to a reduction in trabecular bone density.
While it has been reported previously that gp130 family members, particularly IL-6 and OSM, regulate articu-
ar cartilage breakdown and may play a role in the pathogenesis of various aspects of arthritis (41, 42), gp130 family members have not been studied in growth plate chondrocytes or in longitudinal bone growth. The LIF-Rβ knockout mouse exhibits reduced postnatal skeletal growth (15), but changes in the growth plates of these mice have not been reported. As bone size is normal in IL-6 and IL-11Ra1 knockout mice (16), the reduced bone size observed in gp130STAT1/3 knockout mice may relate to impaired STAT1/3-mediated signaling in response to gp130LIF-Rβ heterodimerization. LIF, acting via LIF-Rβ and the STAT1/3 pathway, may play an essential role in maintaining a high level of chondrocyte proliferation during skeletal growth.

No change in osteoblast or osteoclast activity or trabecular bone volume was detected when gp130-dependent STAT1/3 signaling was abolished in gp130STAT1/3 mutants, suggesting that STAT1/3 is not required for normal bone turnover in the basal state, despite its involvement in the generation of RANKL in response to gp130 cytokines by osteoblastic stromal cells in vitro (43). In contrast, the gp130Y757F/Y757F mutant mice demonstrated high levels of both bone formation and bone resorption, resulting in an osteopenic phenotype. The increased osteoclast number, but not the increased osteoblast number, observed in the gp130Y757F/Y757F mutant mice was reproduced in ex vivo cultures of bone marrow cells from these mice. Here we saw that formation of functional osteoclasts induced by the same dose of RANKL was dramatically elevated in the gp130Y757F/Y757F mice but was not changed in gp130STAT1/3 mice. This result suggests that the defect in osteoclast number observed in vivo may relate to an inherent change in the osteo-
clast precursor population and is not mediated by systemic hormones or altered paracrine signaling from osteoblasts. Since these mice also display increased numbers of macrophage CFU progenitors (44), which themselves are able to give rise to osteoclasts, it appears likely that the altered number of osteoclast progenitors in the bone marrow population used for the ex vivo cultures may cause the high level of bone resorption in vivo. To account for this inherent variation in the pre-
cursor population, we examined the osteoclastogenic potential of a more homogeneous population of osteo-
clast precursors (BMMPs). Even in this preparation, in which osteoclast precursor numbers are equal for both genotypes, the precursors from gp130Y757F/Y757F mutant continued to show a heightened osteoclastogenic response to RANKL. Thus, deletion of the gp130-SHP2/Ras/MAPK pathway gives rise to a greater number of osteoclast precursors, and these precursors have a greater maximum response to RANKL, leading to increased osteoclastogenesis.

As a high level of osteoclastogenesis is also observed in LIF-Rβ- and gp130-/- mice (15, 17, 18), our study has demonstrated a novel role for gp130 in the osteoclast cell lineage in controlling osteoclastogenesis. It sup-
ports a model whereby gp130 family members signal through the gp130-SHP2/ras/MAPK pathway to limit osteoclast formation and indicates that this property is contained within the hematopoietic lineage. Previously, it had been recognized that activation of gp130 on osteoblasts by IL-6 indirectly stimulated osteoclast for-
mation in vitro (9, 32, 43) via activation of STAT3 (34). Although STAT3 activation is upregulated in gp130Y757F/Y757F mice, we know that this IL-6-dependent pathway alone is not responsible for the high osteo-
clastogenesis in these mice, as this is retained in gp130Y757F/Y757F:IL-6-/- mice, and osteoclastogenesis is not altered when the gp130-STAT1/3 pathway is abol-
ished in the gp130STAT1/3 mice. Hence, our data sug-
gest a dual role for gp130 in osteoclastogenesis based on its simultaneous expression on osteoblast and osteoclast precursors (45–47), which may be important in balancing the opposing activities of these two cell types. Although gp130 cytokines stimulate osteoclas-
togenesis through the osteoblast lineage, the dominant effect on bone in vivo is the hematopoietic lineage–spe-
cific effect of the gp130 pathway in inhibiting osteo-
clastogenesis (see Figure 6).

The Ras/MAPK pathway not only functions down-
stream of gp130 but also has been implicated in RANKL, IL-4, and TNF-α stimulation of osteoclasto-
genesis from bone marrow macrophage precursors (48–50). While the Ras/MAPK pathway is often impli-
cated in the stimulation of cell proliferation and dif-
f erentiation, this pathway is also involved in cell senes-
cence, depending both on the manner of its activation and on autocrine/paracrine signaling pathways down-
stream of Ras/MAPK itself (51). It is likely that the down-
stream effectors of the MAPK pathway through which gp130 inhibits osteoclastogenesis described here are different from the stimulatory pathway previously described in osteoclastogenic cell culture systems.

Because IL-6 stimulates both bone formation and resorption, and mutation of the SHP2 binding site enhances IL-6 signal transduction (6), we examined whether increased bone formation and resorption in gp130Y757F/Y757F mice is mediated by altered IL-6 signaling by generating compound gp130Y757F/Y757F:IL-6-/- mice. Whereas IL-6-/- mice have no bone abnormality (16, 36), ObS/BS was rescued to levels of wild-type mice in compound gp130Y757F/Y757F:IL-6-/- mice, while Ocs/BS remained high. This high level of bone resorp-
tion without a compensatory increase in bone forma-
tion explains the very low bone mass in the compound mutant mice. It also shows that IL-6 is not responsible for the increased osteoclast formation and bone resorption in those mice, as suggested by the ex vivo studies discussed above.

Furthermore, since gp130Y757F/Y757F marrow cultures did not exhibit increased osteoblast differentiation or mineralization, the high level of bone formation in gp130Y757F/Y757F mice must require paracrine or endocrine signaling. The rescue of the high ObS/BS in
gp130 deletion (gp130−/−) mice is consistent with a requirement for IL-6 for the enhanced osteoblastic response in gp130−/− mice. Furthermore, this condition does not result from gp130−/− mice, this pathway is clearly not required for normal basal bone formation, since this is normal in gp130−/− STAT3/STAT1 mice. Furthermore, this confirms cell culture studies reporting that while IL-6 stimulation of bone formation can be mediated by either gp130 signaling pathway (34), the gp130-STAT1/3 pathway is the major pathway by which IL-6 stimulates osteoblast differentiation (35). Finally, all of this is consonant with our in vitro result that increased osteoclastogenesis in gp130−/− mice does not result from the high level of bone formation and is independent of the effect of this mutation on osteoblasts.

In conclusion, we have defined three key pathways by which gp130 signaling contributes to bone cell interactions and to the determination of normal bone size and bone density (Figure 6a). First, gp130 cytokines, via the STAT1/3 pathway, play an essential role in stimulating chondrocyte proliferation; in the absence of this pathway, premature growth plate closure and reduced bone size is observed (Figure 6b). Second, the gp130-SHP2/Ras/MAPK pathway plays an essential role in inhibiting osteoclastogenesis; in the absence of this pathway, a high level of osteoclastogenesis leads to trabecular and cortical bone loss (Figure 6c). Finally, we confirmed that IL-6 stimulates osteoblast generation through the gp130-STAT1/3 pathway and showed that this pathway is upregulated and responsible for the high level of bone formation in the absence of the gp130-SHP2/Ras/MAPK pathway (Figure 6c). Therefore, gp130 cytokines regulate bone growth and bone remodeling by discrete activation of three pathways in three different cell lineages within the bone, and these distinct pathways may provide new avenues for improving bone growth and bone density.

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