Repair of Local Bone Erosions and Reversal of Systemic Bone Loss Upon Therapy with Anti-Tumor Necrosis Factor in Combination with Osteoprotegerin or Parathyroid Hormone in Tumor Necrosis Factor-Mediated Arthritis

Kurt Redlich,* Birgit Görtz,* Silvia Hayer,* Jochen Zwerina,* Nicholas Doerr,† Paul Kostenuik,‡ Helga Bergmeister,‡ George Kollias,§ Günter Steiner,*¶ Josef S. Smolen,*¶ and Georg Schett*

From the Department of Internal Medicine III, Division of Rheumatology, and the Institute of Biological Sciences, University of Vienna, Vienna, Austria; the Center of Molecular Medicine, Austrian Academy of Sciences, Vienna, Austria; the Department of Pathology, Amgen, Incorporated, Thousand Oaks, California; and the Molecular Genetics Laboratory, Institute of Immunology, Alexander Fleming Biomedical Sciences Research Center, Vari, Greece

Local bone erosion and systemic bone loss are hallmarks of rheumatoid arthritis and cause progressive disability. Tumor necrosis factor (TNF) is a key mediator of arthritis and acts catabolically on bone by stimulating bone resorption and inhibiting bone formation. We hypothesized that the concerted action of anti-TNF, which reduces inflammation and parathyroid hormone (PTH), which stimulates bone formation, or osteoprotegerin (OPG), which blocks bone resorption and could lead to repair of local bone erosions and reversal of systemic bone loss. To test this, human TNF-transgenic mice with established erosive arthritis and systemic bone loss were treated with PTH, OPG, and anti-TNF, alone or in combination. Local bone erosions almost fully regressed, on combined treatment with anti-TNF and PTH and/or OPG, suggesting repair of inflammatory skeletal lesions. In contrast, OPG and anti-TNF alone led to arrest of bone erosions but did not achieve repair. Treatment with PTH alone had no influence on the progression of bone erosions. Local bone erosions all showed signs of new bone formation such as the presence of osteoblasts, osteoid formation, and mineralization. Furthermore, systemic bone loss was completely reversed on combined treatment and this effect was mediated by osteoblast stimulation and osteoclast blockade. In summary, we conclude that local joint destruction and systemic inflammatory bone loss because of TNF can regress and that repair requires a combined approach by reducing inflammation, blocking bone resorption, or stimulating bone formation. (Am J Pathol 2004, 164:543–555)

Chronic inflammation and bone loss are closely linked. This is impressively illustrated in rheumatoid arthritis (RA), a chronic inflammatory disease of the joints, which is characterized by synovial inflammation and cartilage and bone destruction. Tumor necrosis factor (TNF) is a central inflammatory mediator of RA and its therapeutic inhibition leads to dramatic improvement in signs and symptoms of RA.1–4 In addition, TNF has profound effects on bone: overexpression of TNF is not only involved in local bone erosion, but also induces generalized bone loss.5,6 Therefore, TNF can be considered as an important link between chronic inflammation and bone loss.

At a cellular level, TNF affects bone by its potential to inhibit osteoblast function and its ability to stimulate osteoclast formation.7,8 The role of TNF as a potent stimulator of osteoclasts directed attention to the role of osteoclasts in local bone erosion of arthritis. It has been shown that inflammatory bone erosions in human RA and experimental arthritis contain abundant amounts of osteoclasts.9,10 Furthermore, osteoclasts play an essential role in the development of such lesions, as unequivocally demonstrated in arthritis models, which lack osteoclasts. Thus, mice lacking essential mediators of osteoclastogenesis, such as c-Fos or receptor-activator of nuclear factor-κB ligand (RANKL), develop synovial inflammation, but no bone erosions on induction of arthritis by

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Address reprint requests to Georg Schett, M.D., Department of Internal Medicine III, Division of Rheumatology, University of Vienna, Währinger Gürtel 18-20, A-1090 Vienna, Austria. E-mail: georg.schett@akh-wien.ac.at.
immune complexes or TNF overexpression, respectively.11,12

Therapeutic interventions to slow or arrest local bone erosions are important because joint destruction is a major determinant of functional disability in human RA.13 Therapies that inhibit bone erosions target molecules involved in the differentiation and activation of osteoclasts.3,4,14 Inhibition of TNF and interleukin-1 in human RA by antibodies, soluble receptors, or receptor antagonists is important in currently used clinical examples.2–4,14 and experimental therapy with osteoprotegerin (OPG), a decoy receptor of RANKL, slows or even completely blocks bone erosion in animal models of arthritis.10,15,16

Although there is occasional radiological evidence for healing of erosions, it is unclear, however, whether bone erosions, once they have been formed, can regress and be restored by normal bone.17–19 Such healing may require osteoblasts, which is the cell competent to build up new bone. Blockade of osteoclasts slows or, at best, blocks bone erosion, but may not be sufficient in repair of erosions, which likely requires the synthesis and mineralization of bone matrix. Thus, repair of local bone erosions may require new approaches, such as combination of agents that lead to 1) reduction of the inflammatory process of arthritis that fuels bone erosion, 2) blockade of the resorptive capacity of arthritis by inhibiting osteoclasts, and 3) formation of new bone by stimulation of osteoblasts.

We addressed this question using human tumor necrosis factor transgenic (hTNFtg) mice that develop erosive arthritis.8 Treatment was initiated long after local bone erosions and systemic bone loss had already developed. Following the above hypothesis, we applied a TNF-blocker (for interference with the inflammatory response), OPG (to directly affect osteoclast differentiation and activation), and parathyroid hormone (PTH) to stimulate osteoblast activity as single therapies or in combination.

Materials and Methods

Animals and Treatments

The heterozygous Tg197 transgenic mice have been described previously.5 Briefly, mice were made transgenic for the human TNF-α gene construct with an unmodified 5’promoter region, but allowing deregulated human TNF-α gene expression in vivo. Preparation of the hTNF gene constructs used has been described in detail.5 These mice develop a chronic inflammatory and destructive arthritis within 5 weeks after birth. All mice were bred and maintained in a CBAXC57BL/6 genetic background. Mice were fed a standard diet with water ad libitum. All animal procedures were approved by the local ethical committee.

A total number of 68 hTNFtg mice were assessed for clinical signs of arthritis (see below) up to 10 weeks after birth, ie, 5 weeks after initiation of the arthritic process. At this time point mice were matched according to clinical signs of arthritis into six groups as follows: group 1 (n = 12) was sacrificed at the beginning of treatment (week 10) and served as baseline control. Group 2 (n = 8) was left untreated up to week 14 and served as positive control. Group 3 (n = 8) received 10 mg/kg of OPG intraperitoneally every third day. Group 4 (n = 8) received 80 μg/kg of PTH 5 days per week subcutaneously. Group 5 (n = 8) received 10 mg/kg of infliximab (anti-TNF) intraperitoneally every third day. Groups 6, 7, and 8 (each n = 4) received a combination of OPG + PTH, anti-TNF + OPG, and anti-TNF + PTH, respectively, applied as indicated above. Group 9 (n = 12) received anti-TNF + OPG + PTH (combination). Two independent experiments were performed. All mice were littermate controlled. Treatment was applied between weeks 10 and 14 and mice were then sacrificed. Doses of OPG, PTH, and anti-TNF were based on those successfully applied in previous studies on blocking bone resorption, stimulation of bone formation, and anti-inflammatory activity, respectively.10,20,21

Reagents

Fc-OPG and PTH were kindly provided by Amgen (Thousand Oaks, CA). Fc-OPG is a truncated form of human OPG (amino acids 22 to 194) fused to the Fc domain of human IgG1 and produced in Escherichia coli.22 PTH is the 1 to 34 amino terminal peptide of PTH, which contains all functional properties of the full-length hormone. The chimeric monoclonal anti-TNF antibody (infliximab) was from Centocor (Leiden, The Netherlands).

Cell Culture

Cultures of osteoblasts and osteoclasts were performed as previously described.8,23 In brief, bone marrow was removed from femoral bones by flushing with α-minimal essential medium. For culturing osteoblasts, cells were plated at 103 cells/cm2 and cultivated in α-minimal essential medium supplemented with 10% fetal calf serum, 5 mmol/L glycerophosphate, and 100 μg/ml ascorbic acid. After 10 days, histochemical staining of alkaline phosphatase (AP) was performed. Briefly, cells were fixed with neutral-buffered formalin for 20 minutes at 4°C, incubated with substrate solution at room temperature for 15 minutes, rinsed five times with distilled water, and photographed (F601; Nikon, Tokyo, Japan). AP-substrate was prepared from solution A (8 mg naphthol-AS-TR-phosphate/300 μl N,N’-dimethylformamide; both from Sigma, St. Louis, MO) and solution B (24 mg fast blue BB salt/30 ml of 100 mmol/L Tris-HCl, pH 9.6). Both solutions were mixed, 10 mg of MgCl2 was added, and used immediately after sterile filtration. Enzymatic activity was determined in cell lysates that were solubilized with 0.1% Triton X-100. Aliquots (20 μl) of each sample were incubated with 100 μl of AP substrate buffer (100 mmol/L diethanolamine, 150 mmol/L NaCl, 2 mmol/L MgCl2, p-nitrophenylphosphate at 2.5 μg/ml) for 5 to 30 minutes at room temperature. Total cellular protein was determined using the bicinchoninic method (Pierce Chemical Co.,

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Rockford, IL). AP activity is expressed as U/mg protein with 1 U defined as enzymatic activity that released 1 nmol of p-nitrophenol/minute. Bone nodule formation was assessed after 21 days of culture using alizarin red staining (Sigma).

For osteoclast culture, bone marrow mononuclear cells were isolated by Ficoll gradient centrifugation. Cells were plated and cultivated in a minimal essential medium supplemented with 20 ng/ml of M-CSF (R&D Diagnostics, Minneapolis, MN) and 50 ng/ml RANKL (R&D). Tartrate-resistant acid phosphatase (TRAP) staining was performed after 5 days of culture using a leukocyte acid phosphatase kit (Sigma).

Clinical Assessment of Arthritis

Clinical evaluation was started 6 weeks after birth and performed weekly up to week 14. Arthritis was evaluated in a blinded manner as described previously. Briefly, joint swelling was examined using a clinical score graded from 0 to 3 (0, no swelling; 1, mild-; 2, moderate-; or 3, severe swelling of toes and ankle). In addition, grip strength of each paw was analyzed on a wire of 3 mm in diameter using a score from 0 to −4 (0, normal grip strength; −1, mildly-; −2, moderately-; −3, severely reduced grip strength; −4, no grip strength at all). The last evaluation was performed 14 weeks after birth. Animals were sacrificed by cervical dislocation, the blood was withdrawn by heart puncture and the paws of all animals were dislodged for histology.

Joint Histology

Paws were fixed in 70% ethanol and embedded undecalcified in methylmetacrylate (K-Plast; Medim, Buseck, Germany). After polymerization sections of 3 µm were made on a Jung microtome (Jung, Heidelberg, Germany), deplastinated, and stained by Goldner trichrome, von Kossa, and toluidine blue. Bone erosions were assessed at a magnification of ×200 using a Zeiss microscope (Axioskop 2) and the OsteoMeasure program 2.2 (Osteometrics). The following parameters were measured according to international standards: fraction of bone volume of total sample volume (BV/TV), trabecular number (Tb.N), trabecular thickness (Tb.Th), numbers of osteoclasts per bone perimeter (N. OC/B.Pm), and numbers of osteoblasts per bone perimeter (N. OB/B. Pm). In addition, the width of primary spongiosa (Wi.PS) was assessed.

Bone Histomorphometry

Left tibial bones were fixed in 70% ethanol and embedded undecalcified in methylmetacrylate as described above and stained by Goldner trichrome. Histomorphometry was performed at a magnification of ×200 using a Zeiss microscope (Axioskop 2) and the OsteoMeasure program 2.2 (Osteometrics). The following parameters were measured according to international standards: fraction of bone volume of total sample volume (BV/TV), trabecular number (Tb.N), trabecular thickness (Tb.Th), numbers of osteoclasts per bone perimeter (N. OC/B.Pm), and numbers of osteoblasts per bone perimeter (N. OB/B. Pm). In addition, the width of primary spongiosa (Wi.PS) was assessed.

Densitometry

As previously described, bone mineral density (BMD) was quantified in ethanol-fixed, undecalcified bones by peripheral quantitative computed tomography (pQCT) analysis (XCT-960 mol/L tomograph; Norland Medical Systems, Ft. Atkinson, WI; XMICE 5.2 statistical software; Stratec, Pforzheim, Germany). BMD was measured in the tibia for the total zone (ie, the entire bone), the trabecular (ie, the inner 20% of bone mass), and the cortical zone. For the total and trabecular zones, values from two 1.25-mm sections, located 0.5 mm apart, taken 1.5 mm distal to the proximal articular surface were averaged. In contrast, BMD in the cortical zone of the tibia was measured in a 1.25-mm section located 4.0 mm distal to the articular surface. The total and trabecular regions were differentiated from soft tissue using a separation threshold of 1500, whereas for delineating the cortical zone a threshold of 2000 was used.

Serum Parameters of Bone Metabolism and hTNF Levels

Serum levels of deoxypyridinoline (DPD) crosslinks were measured by enzyme immunoassay (Quidel, San Diego, CA) after previous hydrolysis of serum samples according to the manufacturer’s recommendations. Osteocalcin was measured by immunoradiometric assay (Immunopectics, San Clemente, CA). Serum levels of human TNF were assessed by enzyme immunoassay (R&D).

Results

hTNFtg Mice Show Increased Formation of Osteoclasts but Decreased Formation of Osteoblasts

To address the potential of hTNFtg mice to generate osteoclasts and osteoblasts, we investigated differentiation of bone marrow mononuclear cells into osteoclasts and of bone marrow stromal cells into osteoblasts. Formation of TRAP-positive cells on stimulation with M-CSF and RANKL was increased in hTNFtg mice compared to
wild-type controls (Figure 1; a to c). Furthermore, size of osteoclasts was significantly (P < 0.01) increased in hTNF
tg mice compared to wild-type controls (mean ± SEM, 0.0021 mm
± 0.0001 versus 0.0014 mm
± 0.00007) indicating increased osteoclastogenesis. In contrast, AP production (Figure 1; d to f) and bone nodule
formation (Figure 1; g to i) by bone marrow stromal cells of hTNF
tg mice was decreased, suggesting impaired osteoblast differentiation. These data not only confirmed the basis underlying the destructive process namely osteoclast hyperactivity, but also provide an explanation for the lack of repair process in TNF-mediated arthritis, namely reduced osteoblast activity.

Joint Inflammation Is Only Affected by Anti-TNF but Not OPG or PTH

At baseline (week 10), all hTNF
tg mice had already undergone 5 weeks of active arthritis as seen clinically (Figure 2, a and b) and histologically (Figure 2, c and e). Moreover, local bone erosions containing numerous os-
a) Joint swelling scores over age (weeks)

b) Grip strength scores over age (weeks)

Baseline hTNFtg

c) H&E

d) TRAP
teoclasts were detectable (Figure 2, d and f). At this time point treatment was initiated and continued until week 14. Although hTNFtg control mice, as well as mice treated with OPG or PTH alone (Figure 2, a and b) or together (data not shown) progressed in paw swelling and continued to lose grip strength, mice treated with anti-TNF showed a significantly improved course of disease (Figure 2, a and b). Anti-TNF along with OPG (data not shown) or PTH (data not shown) or OPG+PTH (Figure 2, a and b), was equally effective, suggesting that PTH and OPG, alone or in combination, did not significantly alter the efficacy of anti-TNF on joint inflammation.

**Anti-TNF in Combination with OPG or PTH Leads to Repair of Bone Erosions**

As illustrated before (Figure 2, d and f), histological analyses revealed bone erosions in all hTNFtg mice at baseline, confirming a severely destructive disease state at the onset of treatment. The natural course of disease showed a further increase in bone damage, as evident from mice left untreated from week 10 to week 14 (Figure 3, a, c, and d). This is depicted in Figure 3, c and d, which reveals significant loss of juxtaarticular bone (stained black by van Kossa labeling) between baseline and week 14. Treatment with PTH alone (Figure 3, a and f) did not affect this progression, showing a similar degree of bone erosion as found in untreated mice. In contrast, OPG, anti-TNF, and OPG+PTH led to an arrest of bone erosion, indicated by an amount of bone damage similar to that observed in the baseline control group (Figure 3; a, e, and g). However, mice treated with anti-TNF along with OPG, PTH, or OPG+PTH, respectively, had virtually no bone erosions (Figure 3, a and h). This indicated that bone erosions already present at the onset of therapy at week 10 had regressed and that, therefore, local repair processes had been induced by the therapeutic regimens.

On the basis of our observations that bone erosions can regress, we searched for evidence of bone formation within erosions. As depicted in serial sections (Figure 4), bone erosions showed all signs of new bone formation in vicinity to osteoclast-mediated bone resorption. Figure 4a shows subchondral bone erosion stained with toluidine blue. The area below the articular cartilage, which normally contains subchondral bone, is filled with invasive inflammatory tissue, while articular cartilage shows signs of proteoglycan loss. Goldner-trichrome staining of the same lesion shows that this subchondral bone erosion consists of two subcompartments with different bone metabolism. (Figure 4b). Whereas one compartment (adjacent to cartilage) shows a resorption lacuna with an osteoclast, the other subcompartment shows numerous osteoblasts and osteoid formation, but no osteoclasts. This is further illustrated by von Kossa staining of the same lesion. The area between osteoblasts and mineralized bone is filled by unmineralized osteoid (Figure 4c). In contrast, mineralized bone is directly attached by inflammatory tissue at the area where osteoclasts form a resorption lacuna. Finally, as illustrated by calcein fluorescence, mineralization and thus completion of new bone formation occurs only in areas of bone erosions that are covered by osteoblasts and osteoid, but not next to osteoclast-mediated resorption lacunae (Figure 4d). These observations suggest that eroded bone maintained capacity to recover.

**OPG Reduces Osteoclasts and PTH Increases Osteoblasts and Osteoid Formation in Local Bone Erosions**

To test whether the observed therapeutic effects on bone erosions (Figure 3) are related to changes in local bone turnover, we quantitatively assessed osteoclast and osteoblast numbers as well as osteoid formation by osteoblasts within erosions. Osteoclast numbers increased from baseline in untreated hTNFtg mice, reflecting the progression of erosion size (Figure 5a). In parallel, however, osteoblast numbers also increased, indicating skeletal responsiveness to increased bone resorption (Figure 5b). The numbers of osteoclasts and osteoblasts were low in mice subjected to anti-TNF along with either OPG, PTH, or OPG+PTH, which was because of almost complete regression of bone erosions. Thus, osteoclast and osteoblast numbers were closely linked. To evaluate the potential of osteoblasts for repair of bone erosions, we quantified the amount of osteoid produced by osteoblasts within bone erosions as an indicator for new bone formation (osteoid area per osteoblast). Osteoid production was increased on therapy with PTH and particularly on treatment with anti-TNF+PTH or triple (combination) therapy, which suggests that osteoblast stimulation by PTH effectively leads to new bone formation within arthritic bone erosions (Figure 5c).

**Osteoclast Blockade Is Essential for Reversal of Systemic Bone Loss Mediated by TNF**

We have previously reported that hTNFtg mice, similar to patients with RA, suffer from systemic bone loss.6 To address the question if the local changes described above were accompanied by systemic bone loss we evaluated the effects of the different treatment regimens on prolonged osteoporosis. TNF-blockade, when started at week 10, had no effect on BMD, neither total (Figure...
6a) nor trabecular (Figure 6b), compared to untreated hTNFtg control mice (Figure 6, a and b). Likewise PTH alone or in combination with anti-TNF had no effect. In contrast, a significant increase of BMD was observed with OPG therapy (Figure 6; a, b, and e). However, there was an even more dramatic increase in BMD when OPG was combined anti-TNF, PTH, or anti-TNF/H11001 PTH (Figure 6; a, b, and f). Gain in bone density was because of an increase of trabecular bone, whereas cortical bone remained almost unaffected (Figure 6, b and c). These data together indicate that even longstanding inflammatory bone loss is reversible by osteoclast blockade with OPG, or more effectively, if osteoblasts are stimulated or inflammation is blocked simultaneously. Histological correlates of changes in bone density are depicted in Figure 6, d, e, and f, showing reappearance of trabecular bone in the metaphyses of mice treated with OPG (Figure 6e) and, most prominently with combination therapy (Figure 6f).

**Figure 6.** Local bone erosions show all signs of functional bone turnover. In serial sections of a single subchondral bone erosion (arrowheads) signs of bone resorption as well as new bone formation are detectable. a: Staining with toluidine blue reveals proteoglycan loss, indicated by destaining of the AC, articular cartilage; ST, inflammatory synovial tissue; and BO, erosion of subchondral bone. b: Goldner-trichrome staining reveals multinucleated osteoclasts (OCs) as well as cubic mononuclear osteoblasts (OBs) at the front of erosions. c: Von Kossa staining shows an osteoid seam between osteoblasts and bone surface. d: Calcein labeling indicates a mineralization front underneath the osteoid seam. Original magnifications, ×200.

Combinations of PTH, OPG, and Anti-TNF Increase Trabecular Bone Mass by Reducing the Number of Osteoclasts and Stimulating Osteoblasts

We next specified the morphological basis of changes in bone density in hTNFtg mice. Bone volume increased significantly on treatment of hTNFtg mice with OPG alone or most prominently together with anti-TNF, PTH, or anti-
TNF−PTH (Figure 7a). Similar results were obtained in respect to the width of primary spongiosa and trabecular number (Figure 7, b and c), whereas trabecular thickness only increased in mice treated with combination therapy (Figure 7d). Taken together these data indicate a recovering of bone microarchitecture after OPG. Changes in bone mass were accompanied by decreased numbers of osteoclasts in therapies containing OPG as well as increased osteoblast numbers in therapies, which included PTH (Figure 7, e and f). Thus, combination treatment is highly effective because of synergy of osteoclast blockade with osteoblast stimulation.

Finally, to address whether the effects observed in densitometry and histomorphometry correlate with systemic changes of bone metabolism in hTNFtg mice, we analyzed serum levels of DPD and osteocalcin. Levels of DPD, a marker for bone resorption, were significantly decreased after treatment with anti-TNF alone or in combination with PTH. Treatments containing OPG, however, were even more effective, indicating successful blockade of systemic osteoclast activity (Figure 8a). Conversely, PTH, alone or in combination, significantly increased levels of osteocalcin, a marker of osteoblast activation and bone formation (Figure 8b). In contrast, serum levels of human TNF were not altered by treatments (data not shown). Thus, the effects of OPG and PTH on bone are reflected by profound changes in systemic bone metabolism. The combination of osteoclast blockade with osteoblast stimulation has a positive net effect on bone mineralization, in addition to that on local bone erosions.

Discussion

Local and generalized bone loss represent a serious problem for patients with RA. Local bone erosion is a hallmark of RA patients and is considered as irreversible damage associated with increased morbidity and loss of joint function.13 Generalized bone loss in RA patients is a consequence of high inflammatory disease activity, functional impairment, and corticosteroid use, and leads to an increased risk of osteoporotic fracture.25,26 Repair of bone damage, rather than slowing or at best blocking structural damage should therefore be an ambitious aim of anti-rheumatic therapy. Previous data from in vivo experiments and clinical studies have shown that pure blockade of TNF is not sufficient to repair or even just arrest local and systemic inflammatory bone loss, suggesting that a more profound intervention in the bone remodeling process is necessary to achieve this ambitious goal. This study demonstrates that therapeutic regulation of bone metabolism, as achieved with the combination of PTH or OPG, dramatically improves structural joint damage when combined with an agent that controls the chronic inflammatory process, such as anti-TNF antibody. Moreover, combination therapy also led to a dramatic increase of systemic bone mass in mice suffering from longstanding inflammatory disease and severe osteoporosis, ultimately reversing it to the physiological levels of healthy mice.
The rationale for combining TNF-blockade with osteoclast inhibition or osteoblast stimulation to overcome TNF-mediated bone damage comes from in vitro observations on the effects of TNF on osteoclasts and osteoblasts.7 TNF is a potent stimulator of osteoclast differentiation and activation, which is illustrated by increased formation of osteoclasts from mononuclear precursors.27 The effect is mediated by induction of RANKL expression and a synergistic effect with RANKL on nuclear factor-κB and stress-activated/c-Jun NH2-terminal protein kinase, two signaling pathways essential for osteoclastogenesis.8 With respect to the osteoblast lineage, however, TNF is considered a negative regulator that inhibits osteoblast differentiation and function. Decreased in vitro differentiation of bone marrow stromal cells into osteoblasts supports this idea.7,28 Several mechanisms have been described, which explain the regulatory effect of TNF on osteoblasts. TNF inhibits IGF-1 and the transcription factor Runx-2, which are involved in the differentiation process of osteoblasts.29 In addition, TNF inhibits the synthesis of matrix proteins, such as osteocalcin30 and AP,31,32 which are considered to be indicators for osteoblast function. This is illustrated by TNF-transgenic mice, which show a negative net effect of TNF on bone resulting in local bone erosions and generalized bone loss.5,6 Thus, a combined intervention, which slows bone resorption and induces bone formation, seems to be a highly effective strategy to achieve repair of TNF-induced bone damage.

OPG can be considered to be a feasible means of blocking TNF-mediated bone resorption, because it interferes with RANKL, which is essential for TNF-mediated osteoclastogenesis.33 OPG, given in pharmacological doses, effectively blocks bone resorption and increases systemic bone mass.34 This was because of a predominant effect of OPG on trabecular bone, which exhibits a more active turnover than cortical bone.6 OPG also slows local bone erosion in arthritic joints, indicating that TNF and RANKL induce formation of osteoclasts in the synovial membrane, which ultimately erode subchondral bone.10,15,35 Indeed, the fact that in our model OPG treatment results in a positive net effect on bone, suggests that the primary effect of OPG is an effective blockade of bone resorption. However, an inhibitory effect of OPG on osteoblasts also could be observed, attributing to recent experimental findings, which identified RANKL as stimulator of osteoblasts and supporting observations of blunted bone formation after OPG treatment in laboratory animals and humans.6,34,36 In contrast, PTH primarily stimulates osteoblasts and its effects on osteoclasts are of indirect nature.37 PTH, when given cyclically, stimulates bone formation and leads to increased bone mass.38 Because TNF inhibits PTH-mediated effects on osteoblasts,39 therapeutic substitution of PTH seems to
be a feasible strategy to stimulate bone formation in the presence of TNF.

Most importantly, local bone erosions regressed after TNF blockade combined with osteoclast inhibition or osteoblast stimulation. Local erosions are mediated by osteoclasts derived from mononuclear precursor cells within the inflamed synovial tissue. Current therapies for RA, such as blockade of TNF can slow or, at best, arrest progression of local bone erosion.2,3,40 Experimental blockade of osteoclasts by OPG or bisphosphonates also slows this process, but does not induce the repair of such lesions. So far there has been no convincing evidence of repair mechanisms occurring in local bone erosions.10,15,35

Recently, cells expressing PTH receptor have been described within bone erosions and these cells are attached to bone at the erosion front.41 Because of their localization these cells most likely represent osteoblasts, although PTH receptors are expressed by synovial lining cells and fibroblast-like cells within the synovial pannus as well.42 In this study we demonstrate that local bone erosions are an area of dynamic bone turnover. Apart from resorption lacunae formed by osteoclasts, all signs of bone formation are found within such lesions. These include 1) cells with the typical morphological characteristics of osteoblasts localized between erosion front and bone, 2) osteoid formation next to osteoblast at the interface of pannus and bone, and 3) mineralization of newly formed bone. Although there is the potential of repair mechanisms within local bone erosion, bone formation can obviously not completely com-
pensate the erosive process. This may have two reasons: firstly, factors synthesized by synovial inflammatory tissue such as TNF may hamper osteoblast activity and, secondly, excessive osteoclast activity may prevent the recruitment of osteoblasts and thus remodeling of resorption lacunae. Thus, a combined therapeutic approach, which reduces the inflammatory activity of synovial tissue and blocks either osteoclast activity or stimulates bone formation, is necessary to repair local bone erosions in vivo.

Whether these findings of hTNFtg mice are also valid for human RA remains to be determined. Strengths of this model are 1) its chronic progressive course leading to a high burden of disease, 2) a well-defined pathomechanism based on cytokine overexpression, and 3) an excellent possibility to study the local and systemic skeletal effects of inflammation. Although hTNFtg mice represent an attractive model for human RA its limitations should be considered: especially, the model is not an autoimmune driven model of arthritis and hence does not depend on autoantibodies and T cells. Because human RA exhibits autoimmune features, similar studies in autoimmune arthritis models may thus allow gaining further insight in the mechanisms of bone repair during inflammation. Nevertheless, these data not only describe repair mechanisms of inflammatory bone damage but also highlight differences between local and systemic bone loss. Thus, local bone erosion results from a tight interplay between inflammation and unfavorable bone-remodeling processes. Reversal is only possible if these two mechanisms are targeted simultaneously. In fact, both of them appear equally important because neither pure blockade of inflammation nor even complex interference with bone remodeling is sufficient to achieve regression of local bone erosion. In contrast, systemic bone loss is much more a result of osteoclast-mediated bone resorption, which is supported by the fact that OPG per se increases systemic bone mass and by the high efficacy of all OPG-containing combination therapies.

In summary, our data show that joint destruction and generalized inflammatory bone loss are reversible. Bone repair requires therapeutic intervention, which not only controls inflammation but also shifts the balance of osteoclasts to osteoblasts in favor of the latter. These findings open new perspectives in the treatment of RA by demonstrating that bone damage is of reversible nature rather than end-stage damage associated with loss of joint function.

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