Bisphosphonates reduce biomaterial turnover in healing of critical-size rat femoral defects

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Abstract
Treatment of osteoporotic patients with bisphosphonates (BPs) preserves bone mass and microarchitecture. The high prescription rate of the drugs brings about increases in the numbers of fractures and bone defects requiring surgical interventions in these patients. Currently, critical-size defects are filled with biomaterials and healing is supported with bone morphogenetic proteins (BMP). It is hypothesized that BPs interfere with biomaterial turnover during BMP-supported repair of defects filled with β-tricalcium phosphate (βTCP) ceramics. To test this hypothesis, retired breeder rats were ovariectomized (OVX). After 8 weeks, treatment with alendronate (ALN) commenced. Five weeks later, 6 mm diaphyseal femoral defects were applied and stabilized with locking plates. βTCP cylinders loaded with 1 μg and 10 μg BMP2, 10 μg LS1P, an inhibitor of BMP antagonists and 1 μg BMP2/10 μg LS1P were fitted into the defects. Femora were collected 16 weeks post-implantation. In groups receiving calcium phosphate implants loaded with 10 μg BMP2 and 1 μg BMP2/10 μg LS1P, the volume of bone was increased and βTCP was decreased compared to groups receiving implants with 1 μg BMP2 and 10 μg LS1P. Treatment of animals with ALN caused a decrease in βTCP turnover. The results corroborate the synergistic effects of BMP2 and LS1P on bone augmentation. Administration of ALN caused a reduction in implant turnover, demonstrating the dependence of βTCP removal on osteoclast activity, rather than on chemical solubility. Based on these data, it is suggested that in patients treated with BPs, healing of biomaterial-filled bone defects may be impaired because of the failure to remove the implant and its replacement by authentic bone.

Keywords
bisphosphonate, BMP2, fracture healing, osteoporosis, β-tricalcium phosphate

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Introduction
Each year, worldwide more than 8.9 million osteoporosis-related fractures are counted.¹ Loss of bone mass and deterioration of microarchitecture in post-menopausal women are the primary causes for the increase in fracture risk. Bisphosphonates (BPs) are widely considered as the drugs of choice for the prevention of bone loss in post-menopausal osteoporosis. BP bind to calcium phosphate (CaP) minerals with high and prolonged affinity.²,³ Upon cellular uptake, the drugs block osteoclastic bone resorption, resulting in stabilization of bone mass and preservation of structure.²,³ Administration of alendronate (ALN),
the most frequently prescribed BP, results in a reduction of hip and vertebral fractures.4–9 Because BPs are widely used and elderly patients with osteoporosis are prone to fractures, there has been an interest to understand the effects of BP on bone repair. In vivo studies on fracture healing in rodents reported an increase in callus size and a delay in bone remodelling upon treatment with ALN.10–12 Furthermore, in a number of surgical interventions such as spinal fusion or the repair of large bone defects, bone morphogenetic proteins (BMP), such as BMP2 and BMP7, and CaP-based biomaterials need to be used. To achieve clinical efficacy, however, these growth factors are administered in supraphysiological dosages.13 This may be caused, at least in part, by a reduction in bioavailability of the BMP, since the growth factors induce the expression of antagonists at the repair site.14,15 Indeed, an increase in the expression of BMP antagonists was demonstrated during fracture healing and distraction osteogenesis.16–18 Furthermore, biomaterials need to be removed in a timely manner to allow for efficient replacement by authentic bone. To this purpose, it is essential to ensure that implant removal is governed by cellular processes rather than solely chemical dissolution. To improve the bioavailability of BMP2, L51P, an in vitro engineered BMP2 variant with a leucine to proline substitution at amino acid 51, was developed. This modified protein does not activate BMP type I receptor but has affinity for BMP type II receptors and the BMP antagonists noggin, gremlin and others.19 After demonstrating that L51P restored BMP2-mediated osteoblast differentiation by inhibiting noggin in vitro,20 we and others reported that L51P increased the biological efficacy of BMP2 and that the amount of BMP2 required to induce clinically relevant bone formation in a femoral critical-size segmental defect rat model21 or in a calvaria defect model22,23 could be reduced. Since BP were shown to stabilize bone mass and to maintain bone microarchitecture by reducing osteoclastic resorption and total bone turnover, the administration of BP, in the presence of bioavailable BMP2, has been hypothesized to possibly lead to impaired bone healing.

The aim of the present study was to evaluate the effects of ALN on the healing process of a critical-size defect, filled with CaP ceramics, in the femora of osteoporotic rats in the presence of BMP2 and L51P. The data suggest that healing is impaired by ALN due to the attenuated turnover of the biomaterial.

Methods

Experimental design

This study was approved by the local committee for animal experimentation (Bern Committee for the Control of Animal Experimentation, Bern, Switzerland, permit number BE67/16 to WH). Wistar Crl: W1 (Han) rats (female, retired breeders, 8–10 months old and weighing 285–580 g) were purchased from Charles River (Sulzfeld, Germany). The animals were housed in the Central Animal Facility of the Medical Faculty, University of Bern, Switzerland, in compliance with the Swiss Federal Government guidelines for care and use of experimental animals. Animals were allocated to one of three groups (sham, ovariectomized (OVX)/vehicle and OVX/ALN) based on their bodyweight such that the average weight of the animals was not statistically different among the groups. For this purpose, the animals, after equilibration in the facility and before OVX, were divided into three groups (high, medium and low weight) and the animals from each weight group were randomly allocated to each experimental group using a Random Number Generator for iPad. The experimental design and the time course of the study are depicted in Figure 1. Briefly, at \( t = 0 \), animals were either OVX or sham-operated. ALN (OVX only) or vehicle (sham/OVX) treatment was initiated 8 weeks after OVX and continued until sacrifice. A critical-size defect was applied in the left femur 5 weeks after the onset of vehicle/ALN treatment (13 weeks post-OVX) and stabilized with a rigid osteosynthesis system (RatFix™, RISystem AG, Davos, Switzerland). For all surgical procedures, the rats were anaesthetized by subcutaneous injections (1 ml/kg body weight) of a 1:2 mixture...
of ketamine hydrochloride (33 mg/kg body weight; 100 mg/ml) and xylazine hydrochloride (13.3 mg/kg body weight; 20 mg/ml). During surgery, the animals were placed on a heating pad to prevent hypothermia. Group sizes were \( n = 6 \). Sixteen weeks after femoral surgery, euthanization of animals was performed using \( \text{CO}_2 \), followed by cervical translocation, and samples were collected for micro-computer tomography (microCT) and histological analysis. In total, 12 experimental groups were used, encompassing a total of 72 rats.

**Ovariectomy**

After shaving and disinfection, the ovaries were approached through two 2.5 cm flank incisions at the mid-dorsum. The skin was detached from the underlying muscles before incising the muscles. The ovaries were identified, clamped and removed (gently pulled through the incisions and a haemostat was placed between the oviduct and the ovaries). The oviducts were ligated and a cut was made between the haemostat and the ovaries. Haemostasis was controlled before replacing the ligated oviducts in the abdomen. The muscles were identified, clamped and removed (gently pulled through the incisions and a haemostat was placed between the oviduct and the ovaries). The oviducts were ligated and a cut was made between the haemostat and the ovaries. Haemostasis was controlled before replacing the ligated oviducts in the abdomen. The muscle layer was closed with absorbable sutures, and the skin was sutured with non-absorbable thread. *Sham*-operated animals underwent the same surgical procedures, except for the ligation of the oviducts and the removal of the ovaries.

**BP treatment**

ALN solution was prepared as previously described\(^{24} \) and dissolved in 0.9% NaCl. ALN (1.61 \( \mu \)mol/kg body weight; 2 ml/kg bodyweight) or vehicle (0.9% NaCl solution; 2 ml/ kg body weight) was subcutaneously injected 8 weeks after *OVX* twice weekly until sacrifice. After 8 weeks, bone mass was significantly reduced in *OVX* animals as compared to *sham* controls.

**Loading of ceramic \( \beta \)-tricalcium phosphate cylinders with peptides and release kinetics**

Ceramic \( \beta \)-tricalcium phosphate (\( \beta \)TCP) cylinders (diameter 5 mm, length 6 mm and porosity 75%) were used as described previously\(^ {21} \) and were generously provided by the Robert Mathys Foundation (Bettlach, Switzerland). The BMP2 and L51P proteins were expressed in *Escherichia coli* and were kindly provided by Prof. W Sebold, University of Würzburg, Germany.\(^ {19, 25, 26} \) For the measurement of the in vitro release of L51P and BMP2 from the ceramics, 1 or 10 \( \mu \)g of BMP2 or L51P was dissolved in 25 \( \mu \)l of deionized water and adsorbed onto \( \beta \)TCP carriers (five groups, \( n = 6 \) per group), as recommended by the manufacturer (https://vimeo.com/130984695). Thirteen weeks after *OVX* and five weeks after the commencement of the ALN treatment, the central part of the left mid-diaphyseal femur (6 mm) was surgically removed. After shaving and disinfection of the skin, a longitudinal incision in line with the left femur was cut on the lateral thigh. The interval between the aster lateralis and the biceps femoris was developed to expose the bone, the *gluteus superficialis* tendon was detached from the trochanter tertius and the RatFix™ System (RISystem AG, Davos, Switzerland) was mounted onto the intact femora using six interlocking screws. Subsequently, using two Gigli saws, two osteotomies were created between the two central screws, and the central mid-diaphyseal bone fragment was collected for further analysis. Care was taken not to harm the surrounding periosteum. Debris was removed by rinsing the defect site with sterile physiological saline solution. Subsequently, a \( \beta \)TCP implant, which was stabilized with a non-absorbable polypropylene thread, was inserted into the defect and the wound was closed. After the animals recovered from anaesthesia, analgesia was performed for 3 days, using Temgesic (0.0075 mg/kg body weight; 0.3 mg/ml), and full load bearing and unrestricted cage activity were allowed. Movement or instability of the implants and failure of the osteosynthesis were monitored using high-resolution X-ray imaging immediately after surgery. After 16 weeks, the implants, together with the surrounding tissues, were harvested and analysed.

### Table 1. BMP2 and L51P loading of \( \beta \)TCP carriers.

<table>
<thead>
<tr>
<th>Experimental groups</th>
<th>I</th>
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<tr>
<td></td>
<td>Unloaded</td>
<td>1 ( \mu )g BMP2</td>
<td>10 ( \mu )g BMP2</td>
<td>10 ( \mu )g L51P</td>
<td>1 ( \mu )g BMP2/10 ( \mu )g L51P</td>
</tr>
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\( \beta \)TCP: \( \beta \)-tricalcium phosphate; BMP: bone morphogenetic protein.
X-ray

After surgery, the integrity of the surgical site was documented using high-resolution radiography, applying the following image acquisition parameters: 25 kV and 10 s acquisition time (MX-20, Faxitron X-Ray Corporation, Edimex, Le Plessis, France). No animals needed to be excluded from the experiment post-operatively because of incorrect defect localization and/or plate positioning.

Peripheral quantitative computed tomography

The effects of OVX on bone mass and structure were evaluated using in vivo peripheral quantitative computed tomography (pQCT; XCT Research SA, Stratec Medizintechnik GmbH, Birkenfeld, Germany) measurements at the left distal femur and proximal tibia, 7 weeks after OVX, using the following parameters: voxel size, 70 μm; High Voltage, 50 kV; and acquisition time, 10 min. For pQCT analysis, animals were anaesthetized as described above.

Micro-computer tomography

The tissues for histological and microCT analysis were fixed with 4% paraformaldehyde in Phosphate Buffered Saline (PBS) for 24 h, and subsequently transferred to 70% ethanol. Polyether ether ketone (PEEK) plate and Titanium screws remained in situ for these analyses. For microCT analysis (MicroCT40, SCANCO Medical AG, Brüttisellen, Switzerland), the built-in software from SCANCO was used (SCANCO Module 64-bit; V5.15). The long axis of the femur was oriented orthogonally to the axis of the X-ray beam. The X-ray tube was operated at 70 kVp and 114 μA, and the integration time was set at 200 ms. The measurements were performed perpendicular to the longitudinal axis of the femora. Bone growth was evaluated in the area between the two central screws of the fixation systems. To distinguish between soft and mineralized tissues, the tissue was segmented into two tissue types based on their greyscale (grey-level coded mineralization density), that is, <200 Hounsfeld unit (HU) for soft tissues and >200 HU for mineralized tissues (βTCP carrier, mineralized cartilaginous callus, woven and lamellar bone).27 Cortical thickness was measured using the SCANCO built-in algorithm. The analysis was performed with a voxel size of 8 μm.28

Histological analysis and histomorphometry

After microCT analysis, the tissues were embedded in methyl methacrylate as described previously.29 Thereafter, ground sections of approximately 200 μm were prepared (Leica SP1600, Leica Microsystems, Glatthugg, Switzerland). The sections were polished and stained with McNeal tetrachrome.30,31 Microphotographs were taken using a Nikon Eclipse E800 microscope (Nikon Inc., Egg, Switzerland). Bone formation and implant turnover were determined by the ImageJ trainable segmentation plugin; Waikato Environment for Knowledge Analysis (WEKA) automated segmentation histomorphometry32 on 4 serial McNeal tetrachrome-stained ground sections per implant. The sum of the surface area of bone and implant was determined for each animal (four sections/animal). The total bone surface area and implant surface area per treatment group were computed.

Uterus dry weight

Uteri were collected post-mortem, stored overnight at room temperature, and the dry weight was measured 24 h after collection.

Statistical analysis

The following statistical analyses were performed using GraphPad Prism 7 for Windows (GraphPad Software, San Diego, California, USA, www.graphpad.com). Unpaired t-test was used for bodyweight and uterus dry weight analysis; two-way analysis of variance (ANOVA) with Tukey post hoc was applied to pQCT, microCT and histomorphometry analysis. Mean values with standard deviations are shown. Significant at p < 0.05. For multiple testing, p_adj < 0.05 was considered significant.

Results

Release of BMP2 and L51P from βTCP ceramics

BMP2 and L51P were released in vitro with identical kinetics over the course of 16 days, as determined by ELISA (data not shown).

Bone loss in OVX rats and the effects of BP

The effect of OVX on bone mass was determined in vivo by pQCT analysis in proximal tibiae and distal femora at time of OVX and 7 weeks thereafter. Cortical and trabecular densities did not vary between OVX and sham groups at the time of surgery. After 7 weeks, both cortical and trabecular densities were significantly decreased in OVX animals in measurements at four different tibial sites (tibia position 1: 1122.4 ± 49.4 mg/cm^3 vs. 1022.6 ± 42.3 mg/cm^3, p < 0.0001; 248.2 ± 60 mg/cm^3 vs. 203 ± 57.3 mg/cm^3, p = 0.0002; Figure 2).

To confirm the efficiency of OVX, uterine dry weight was measured post-mortem. Uterine dry weight was reduced by 40% in OVX animals as compared to that in sham-operated animals. (sham vs. OVX + vehicle, 248.4 ± 62.9 mg vs. 138.7 ± 41.4 mg, p = 0.0001; sham vs. OVX + ALN, 248.4 ± 62.9 mg vs. 143.2 ± 37.3 mg, p = 0.0001; Online Supplemental Figure 1).
Treatment with ALN leads to an increase in cortical thickness

To assess the effects of ALN treatment in OVX animals, cortical thickness was determined by microCT analysis of the mid-diaphyseal fragments collected during the creation of femoral critical-size defects (corresponding to 13 weeks after OVX and 5 weeks after onset of the ALN treatment). Administration of ALN resulted in a significant increase of the cortical thickness as compared to that in vehicle treated OVX animals (0.64 ± 0.06 mm vs. 0.58 ± 0.05 mm, p = 0.0066; Figure 3).

Bone formation is stimulated by BMP2 and implant turnover is reduced by ALN

Sixteen weeks after implantation, no bone formation and osseointegration of the CaP implant was observed in the critical-size defects upon insertion of unloaded βTCP (control) cylinders irrespective of sham/OVX and vehicle/ALN treatments (Figures 4(a), (f), (k) and 5(a)). Similarly, no bone formation was found when the defects were filled with 10 mg L51P-loaded βTCP cylinders, again independently of the applied treatment protocols (sham, 1.55 ± 0.39 mm² (unloaded) and 1.84 ± 0.31 mm² (L51P), p = 0.9998; OVX/vehicle, 1.37 ± 0.27 mm² and 2.14 ± 0.81 mm², p = 0.9942; OVX/ALN, 1.87 ± 0.57 mm² and 2.89 ± 2.14 mm², p = 0.9708; Figures 4(d), (i), (n) and 5(a)). Some bone formation was detected in defects filled with 1 mg BMP2-loaded βTCP cylinders (Figure 4(b), (g), (l)), but quantitative histomorphometry revealed no significant increase in bone formation as compared to unloaded βTCP cylinders, irrespective of sham/OVX and vehicle/ALN treatments (sham, 1.55 ± 0.39 mm² vs. 3.12 ± 2.62 mm², padj = 0.9272; OVX/vehicle, 1.37 ± 0.27 mm² vs. 4.25 ± 2.82 mm², p = 0.6185; OVX + ALN, 1.87 ± 0.57 mm² vs. 5.76 ± 4.24 mm², p = 0.2954; Figure 5(a)). Induction of bone formation was observed with βTCP cylinders loaded either with 10 μg BMP2 (sham, 1.55 ± 0.39 mm² vs. 12.41 ± 5.02 mm², p = 0.0003; OVX/vehicle, 1.37 ± 0.27 mm² vs. 10.41 ± 2.37 mm², p = 0.0004; OVX/ALN, 1.87 ± 0.57 mm² vs. 14.49 ± 4.06 mm², p = 0.0001; Figures 4(e), (h), (m) and 5(a)) or 1 μg BMP2/10 μg L51P (sham, 1.55 ± 0.39 mm² vs. 9.86 ± 4.51 mm², p = 0.0031; OVX/ALN, 1.87 ± 0.57 mm² vs. 12.62 ± 5.95 mm², p = 0.0001; Figures 4(e), (o) and 5(a)). In OVX
animals receiving βTCP cylinders loaded with 1 μg BMP2/10 μg L51P (Figure 4(j)), no significant increase in bone formation was detected (1.37 ± 0.27 mm² vs. 5.81 ± 2.84 mm², p = 0.1356). As previously observed, bone formation was associated with implant turnover in sham-operated animals. An increased volume of residual ceramic material remained in the defect site in ALN-treated animals as compared to vehicle-treated animals receiving βTCP implants with identical protein loads (10 μg BMP2 (OVX/vehicle vs. OVX/ALN, 1.99 ± 0.52 mm² vs. 5.48 ± 1.62 mm², p_adj < 0.0001; Figures 4(m) and 5(b)); 1 μg BMP2/10 μg L51P (OVX/vehicle vs. OVX/ALN, 1.39 ± 0.72 mm² vs. 4.33 ± 1.20 mm², p_adj < 0.0001; Figures 4(o) and 5(b)). Although minimal extra-implant/peripheral bone formation was observed in vehicle-treated animals whose defects were filled with 10 μg BMP2 and 1 μg BMP2/10 μg L51P loaded βTCP cylinders (Figure 4(c), (e), (h), (j)), respectively, extensive peripheral bone formation was observed in defects from ALN-treated animals (Figure 4(m), 4(o)) with identical βTCP loading conditions. After 16 weeks, microCT renderings of the defect sites revealed that βTCP cylinders formed an interface with the host bone. In defects from vehicle-treated animals in which implants were loaded with 10 μg BMP2 or 1 μg BMP2/10 μg L51P, bone formation
occurred from either the distal or proximal end of the defect and did not fill the entire implant. In contrast, defects in ALN-treated animals filled with \( \beta \)-TCP cylinders loaded with 10 mg BMP2 or 1 mg BMP2/10 mg L51P were characterized by a bone continuum throughout the entire \( \beta \)-TCP implant (Figure 6). MicroCT analysis of serial cross-sections along the longitudinal axis of the defects suggested that the surface area of bone was decreasing from the bone–implant interface towards the centre of the implant. Evaluation of cross-sectional 2-D microCT images confirmed the reduction of the bone surface area from the bone–implant interface towards the centre of the defect (Figure 7).

Discussion

BP are the most frequently prescribed drugs to preserve bone mass and microarchitecture. Treatment with BP and the consequential maintenance of the bone substance results in a decrease in the risk to sustain wrist, vertebral and hip fractures in patients suffering from osteoporosis.\(^4\)\(^,\)\(^7\)\(^,\)\(^9\) Because prolonged treatment with BP causes bone turnover to slow down, and because the concept of a drug holiday during BP treatment is controversially discussed,\(^33\) it is an open question whether bone repair is impaired by concurring treatment with BP in osteoporotic patients. The present study aimed to answer several aspects of repair of a critical-size femoral defect, with \( \beta \)-TCP as a bone filler, in estrogen deficient osteoporotic rats treated with BP such as (i) the preservation of the previously described synergistic effect of BMP2 and L51P on bone healing, (ii) the effects of BP treatment on bone repair and remodelling of the defect site and (iii) the turnover of a \( \beta \)-TCP implant under conditions of blocked osteoclast activity. The data demonstrate that, as was previously shown,\(^21\) L51P causes an increase in the effects of BMP2 on bone formation in sham-operated animals, leading to an increase in bone volume at the defect sites 16 weeks post-implantation compared to animals with implants loaded with 1 mg BMP2 only or unloaded controls. L51P alone, at 10 mg, did not induce any detectable formation of bone. In animals treated with ALN, bone formation was induced in defects filled with cylinders loaded with 10 mg BMP2 and with 1 mg BMP2/10 mg L51P, respectively.

One common feature observed in vehicle-treated control animals, which received implants loaded with either 10 mg BMP2 or 1 mg BMP2/10 mg L51P, is the presence of intact \( \beta \)-TCP ceramic material surrounding the newly formed bone in a ring-like structure. Since the diameter of the \( \beta \)-TCP implants exceeds the diameter of the femoral diaphysis, we hypothesized that with the present experimental setup, bone formation is constrained to a volume similar to the original bone volume. It would be of particular interest to investigate whether the remaining implant will be filled with bone and remodelled or removed without additional bone formation, if the period allowed for repair is prolonged.

Compared to the newly formed bone in the critical-size defects from control animals, bone formation in animals receiving both ALN and 10 mg BMP2 or 10 mg L51P/1 mg BMP2 exceeded the original bone volume and the
perimeter of the implant. Besides the excessive bone formation, implant turnover was greatly reduced in animals treated with ALN. Quantitative evaluation of the remaining βTCP ceramics revealed that the volume of βTCP in ALN-treated animals was doubled when compared with the volume of βTCP in vehicle-treated animals upon loading of the ceramics with 10 μg BMP2 and 10 μg L51P/1 μg BMP2, respectively. As reported previously, bone formation and growth into the porous βTCP implants are prerequisites for the turnover of the ceramics. The inhibition of osteoclastic bone resorption by ALN, however, decreased the dissolution of βTCP ceramics, demonstrating the implant turnover to depend on osteoclast activity and not on the ceramic’s chemical solubility.

In physiological bone remodelling, coupling of osteoclast and osteoblast activities is crucial for the maintenance of mass and structure of bone. In the presence of supraphysiological levels of BMP2, bone formation and resorption are uncoupled. As a consequence, in our study, bone formation induced by BMP2 was not affected in animals treated with ALN, despite the block of resorption. When animals are treated with ALN, irrespective of BMP2 levels, bone remodelling will not take place and therefore primary woven bone will not be replaced by secondary, mechanically more competent lamellar bone.

The present study provided evidence on two crucial aspects relevant to reconstructive orthopaedic surgery. Firstly, it corroborated the usefulness of the strategy to block endogenous BMP antagonists to improve the bioefficacy of exogenously added BMP. Secondly, it demonstrated the necessity for functional cellular resorption for the removal of βTCP ceramics from a repair site. There are, however, also limitations of the present model. The duration of the study was too short for a complete removal of the βTCP biomaterial. Furthermore, no biomechanical testing of the partially or fully healed defects could be performed. Despite these limitations, however, the data suggest that in patients treated with BP, healing of βTCP-filled bone defect may be impaired because of the failure to

Figure 6. MicroCT 3D renderings of femoral defects. Representative renderings from critical-size femoral defects filled with cylinders loaded with 10 μg BMP2 are depicted for (a) sham, (b) OVX and (c) OVX/ALN. Renderings from defects filled with ceramics loaded with 1 μg BMP2/10 μg L51P are shown in panels (d) sham, (e) OVX and (f) OVX/ALN. Bone formation occurs preferentially at the proximal and distal ends of the implants in sham and OVX control animals irrespective of BMP2 and BMP2/L51P loading of the implants (a, b, d, e). In animals treated with ALN, bone formed from both the distal and proximal ends of the defects, the newly formed bone bridging the defect irrespective of BMP2 and BMP2/L51P loading of the implants (c, f). ALN: alendronate; OVX: ovariectomized; BMP: bone morphogenetic protein; microCT: micro-computer tomography.
remove the implant material and its replacement by authentic bone.

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**Supplemental material**
Supplemental material for this article is available online.

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