Prophylactic effects of propranolol versus standard therapy on a new model of disuse osteoporosis in rats

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Running title: Propranolol prevent immobilization-induced bone loss

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Abstract

Disuse by bed rest, limb immobilization or space flight causes rapid bone loss by arresting bone formation and accelerating bone resorption. Propranolol (a non-selective β-adrenergic antagonist) has been shown to improve bone properties by increasing bone formation and decreasing bone resorption in an ovariectomy-induced rat model. However, no studies have compared the osteoprotective properties of propranolol with well accepted therapeutic interventions for the treatment and prevention of immobilization/disuse osteoporosis. To clarify this, we investigated the effects of propranolol compared with zoledronic acid and alfacalcidol in a new animal model of immobilization/disuse osteoporosis. Three month old male Wistar rats were divided into five groups with six animals in each group: (1) immobilized (IMM) control; (2) normal control; (3) IMM + zoledronic acid (50µg/kg, intravenous single dose); (4) Imm + alfacalcidol (0.5 µg/kg, per oral daily); (5) IMM + propranolol (0.1mg/kg, subcutaneously 5 days/week) for 10 weeks. In groups 1 and 3-5, the right hind-limb was immobilized. At the end of treatment femurs were removed and tested for bone porosity, bone mechanical properties and cortical micro-architecture. Treatment with propranolol induced greater reductions in the bone porosity of the right femur and improved mechanical properties of the femoral mid-shaft femur in comparison to IMM control. Moreover, treatment with propranolol also improved microarchitecture of cortical bones when compared with IMM control, as indicated by scanning electron microscopy. The anti-osteoporotic property of propranolol was comparable with zoledronic acid and alfacalcidol. This study shows that the bone resorption induced by immobilization/disuse in rats can be suppressed by treatment with propranolol.

Keywords: Propranolol. Immobilization. Rat model. Osteoporosis. Bone strength
Introduction

Osteoporosis is described as a multifactorial disease characterized by low bone mineral mass and deteriorated microarchitecture of the bone tissue leading to the enhanced bone fragility and increase in the risk of debilitating fractures. Immobilization is one of the important causes of osteoporosis. Disuse (unloading) osteoporosis occurs in patients with spinal cord injuries, patients confined to prolonged bed rest, and astronauts exposed to microgravity during space flight. Disuse osteoporosis also occurs in areas of low bone stress around orthopaedic implants and from limb immobilization after surgery [1]. The main clinical manifestation of long-term or short-term immobilization is represented by increased fracture liability [2]. The main feature during immobilization is a dramatic increase in bone resorption and a decrease in the bone formation [3]. Inspite of continuous and aggressive treatment, immobilization requires a very long time for bone to recover its bone mineral density and mechanical strength [4]. As the goal of today’s medicine shifts more towards disease prevention rather than treatment, prophylactic therapy seems to be the promising choice for disuse osteoporosis.

Zoledronic acid (ZOL), a nitrogen-containing bisphosphonate, is a potent anti-resorptive agent which have been approved and in use for treatment of various disorders characterized by increased osteoclast-mediated bone resorption due to, for example, estrogen depletion or aging/disuse. ZOL interferes with osteoclastic activity by preventing osteoclast formation and osteoclast-bone resorptive activities and by inducing osteoclast apoptotic cell death [5]. Alfacalcidol (ALF) is a prodrug of active vitamin D3, a calcium-regulating hormone which is widely accepted as a baseline treatment for osteoporosis. Supplementation with vitamin D increases muscle strength and thus may reduce the risk of fractures [6]. It has been demonstrated
that the administration of vitamin D analogs diminished the effect of immobilization in the development of osteoporosis [7]. Previously, we studied the preventive therapy of administering ALF to rats immediately after ovariectomy and found that ALF sustained or increased bone mass by suppressing bone resorption while maintaining bone formation [8, 9].

Propranolol (PRO), a non-selective $\beta$-adrenergic antagonist, has been shown to improve bone properties in different experimental models of bone disorders [8–13]. Results of some prior epidemiological studies confirm the hypothesis that $\beta$-blockers use is associated with a decrease in fracture risk [14-16]. According to the available evidence, $\beta$-adrenergic pathway of sympathetic nervous system is a major transmitter pathway that mediates unloading-induced bone loss through suppression of bone formation by osteoblasts and enhancement of resorption by osteoclasts. Studies have demonstrated that PRO could be used to prevent the induced mechanical unloading bone loss [17, 18]. Moreover, studies have demonstrated that low doses of PRO suppress bone resorption by inhibiting receptor activator of nuclear factor kappa-B ligand (RANKL) mediated osteoclastogenesis as well as inflammatory markers without affecting haemodynamic parameters [13]. This result is supported by a previous finding, which showed that PRO stimulates osteoprotegerin (OPG) on its own in osteoblast cells [17]. PRO have been recommended as one of the first-line treatment drugs for hypertension and have been widely used in cardiovascular disease. Based on the available data, nearly 45 % of older population suffers from cardiovascular disease and osteoporosis. Therefore, the advantage of a dual-benefit effect of only one treatment on both heart and skeletal systems is of great interest [9].
With this possibility in mind, the use of PRO was investigated which may lead to a better treatment for immobilization osteoporosis when compared to well accepted therapies such as ZOL and ALF in a rat model of immobilization osteoporosis. A new immobilization model is reported in this study which was very efficient in inducing significant long term hind-limb disuse.

**Materials and methods**

**Drugs, chemicals and other materials**

ZOL was obtained from Naprod Life Sciences, Maharashtra, India. PRO, ALF, xylene and ether was obtained from Aurobindo Pharma (Hyderabad, India), Glaxo Smithkline Pharmaceuticals (Mumbai, India) and S.D. Fine chemicals (Mumbai, India), respectively. Ketamine and xylazine was obtained from Neon Pharma, Mumbai and Indian Immunologicals Ltd., Hyderabad, respectively.

**Pre-clinical study design**

Three months old male Wistar rats of weighing 170–180g were included for the study. Animals were maintained under controlled temperature at 25°C ± 2°C with 12hr light/dark cycle with food and water and provided ad libitum. The experiments were conducted as per the CPCSEA (Committee for the Purpose of Control and Supervision of Experiments on Animals) guidelines after obtaining ethical clearance from the Institutional Animal Ethical Committee.
Description of a new rat model for disuse osteoporosis

The framework for right hind-limb immobilization of rats was prepared with a durable and high quality PVC (polyvinyl chloride) coated welded iron mesh. The mesh was 16 cm long and 6 cm wide with a pore size of 1.8 mm, as shown in fig. 1A. PVC coated iron mesh was then wrapped with microporous adhesive surgical tape as shown in fig. 1B, to prevent injury to the animal’s body. The prepared framework can be easily fitted and fixed to a Wistar rat with body weight of 160-180 grams. Firstly, the reasons for using PVC coated welded mesh iron mesh were based on cost effectiveness, light weight, softness, suitability for sterilization and biocompatibility. Second and most interestingly, the advantage of this framework was to prevent the animal from destroying the inner microporous adhesive tape, which was wrapped to the animal’s abdomen for immobilizing the hind-limb. To maintain retention of the framework on the animals, 2-3 layers of microporous adhesive surgical tape (5 cm wide) was placed around the animal’s abdomen. Microporous adhesive tape usually have a hypoallergenic adhesive which is designed to hold firmly onto skin and underlying layers of tape, but to remove easily without damaging the skin. Moreover, it allows air to reach the skin. It helps to protect the skin from irritation, abrasion and infection.

All animals were housed singly in cages for 2 weeks for acclimatization to vivarium conditions. Before immobilization of the right hind-limb, rats were anesthetized with a combination of ketamine (80 mg/kg) and xylazine (10 mg/kg), intraperitoneally. Each animal’s lower torso and right hind-limb were trimmed of all hair using an automatic hair trimmer (Fig. 1C & 1D). A protective barrier wipe was applied to the lower right hind-limb, as shown in fig. 1E. It provides a barrier film layer on skin used under microporous adhesive tape, to help protect against
irritation, excoriation and adhesive build-up. The right hind-limb was immobilized against the abdomen with the hip joint in flexion and tibiofemoral joint in extension, using two layers of microporous adhesive surgical tape (4 cm wide, as shown in fig. 1F. The framework was encased against the abdomen using 6 cm microporous adhesive tape. It is a necessary step to keep the hind-limb in a fixed immobilized position for longer duration (Fig. 1G). Within 24 h, the rats were able to walk on three legs with no obvious discomfort in locomotion or feeding inside the cage. Throughout the 10 week treatment period the animals were checked daily. Moreover, there was no need to re-adjust the framework as it was able to immobilize the animal successfully for a longer duration, as compared to the conventional methods with no indication of skin ulceration, oedema, sores, or swelling.
Fig. 1. Pictorial representation of rat hind-limb immobilization method. (A) The framework is made of PVC coated iron mesh (measurements in cm), (B) to prevent injury to the animal’s body, PVC coated iron mesh was covered with microporous adhesive tape (measurements in cm), (C) rat’s lower torso and right hind-limb were trimmed of all hair using an automatic hair trimmer, (D) black circle depicts the trimmed area, (E) protective barrier wipe was applied to the lower right hind-limb (arrow), (F) right hind-limb was immobilized using two layers of microporous adhesive tape, (G) framework encased against the abdomen using microporous adhesive tape (measurements in cm).
Experimental procedure

The rats were divided into 5 equal groups (n=6). The non-immobilized control group and the immobilized (IMM) control group, served as negative and positive controls, respectively. Normal (non-immobilized) control and IMM groups were subcutaneously administered vehicle (normal saline, 5 days per week) for 10 weeks. One IMM group was treated with single intravenous dose of ZOL with 50 µg/kg, administered into tail vein as a slow intravenous injection over 30 s under light inhalation anaesthesia. One IMM group was orally administered 0.5 µg/kg of ALF daily for 10 weeks. Treatment on the remaining IMM group was initiated with PRO at a dose of 0.1 mg/kg, injected subcutaneously 5 days per week, for 10 weeks. The medication dosages used in this experiment were selected from previous studies on rat osteoporosis model [8, 9]. The dose of ZOL was selected based on the dose response study carried out in our laboratory. After 10 weeks of drugs or vehicle administration the animals were humanely sacrificed under anesthesia and the various bone parameters were evaluated.

Measurement of femoral porosity

The femurs of all animals were scanned with foX-Rayzor, which is a portable X-ray inspection system equipped with “Calculate histogram” tool software, according to the method described by Khajuria et al [9]. Briefly, for X-ray analysis, whole femur was divided into four equal fields, which includes distal femoral epiphysis (R1), femoral shaft (R2 and R3) and proximal femur (R4).
Bone mechanical tests

Femur strength was assessed by three-point bending as previously described [9]. Briefly, femurs were removed from the -20 ºC freezer and rehydrated in a saline solution for 4 h at room temperature. Hydrated weight of the bones was determined using a four decimal place digital scale. The length of the femurs was measured by using a caliper to determine possible effect on skeleton growth. Specimens were placed on two supports that were separated by a distance of 12 mm and bent until fracture by lowering the crosshead positioned at the mid-shaft at a constant speed of 0.033 mm/s. From the load-displacement curve, the peak load (N), the ultimate stiffness (N/mm), and the toughness (mJ) were obtained. Ultimate stress (strength) and Young’s modulus were derived from load-deformation curves obtained by using equations described by Khajuria et al [9].

Cortical scanning electron microscopy

After mechanical strength tests, fractured surfaces of right femur were examined by SEM (scanning electron microscopy; Ultra 55, Karl Zeiss Microscopy, Germany) at magnification of X600. A proximal-diaphysis section of the right femur of all rats in each group was rendered anorganic by a 5% sodium hypochlorite treatment. The sections were then rinsed in water, dehydrated in acetone, and dried. SEM examinations were carried out for bone pores [19]. Representative SEM photomicrographs were analyzed for number and size of pores using image analysis software (Sigma Scan Pro software).
Statistical analysis

All data were expressed as the mean ± S.D (standard deviation). For all the data, comparisons between different treatments were analyzed by one-way ANOVA followed by Tukey’s multiple comparison tests. In all cases, a probability error of less than 0.05 was selected as the criterion for statistical significance. Graphs were drawn using Graph Pad Prism (version 5.0 for Windows).

Results

Final body weight

The normal control group had significantly higher body weights than the IMM control group and all treatment groups. There were no statistically significant differences in the weights observed between any of the active treatment groups and that of the IMM control group (Fig. 2).

Fig. 2. Final body weight of the animals. Data are expressed as the mean ± S.D (n=6), evaluated by one-way ANOVA followed by Tukey’s multiple comparison test. *p < 0.05, compared to IMM control group.
**Femoral Length**

The length of the immobilized femurs was not significantly different from that of the non-immobilized femurs of the same rats in the control group. The administration of all therapeutic interventions did not cause any significant change in the length of the immobilized and non-immobilized femurs (data not shown).

**Effect of different treatments on bone porosity**

*Immobilized (right) leg*

The effects of immobilization and subsequent treatment with ZOL, ALF and PRO on the porosity of the right femur were measured by X-ray imaging is shown in fig. 3 (a–d). X-ray transmission intensity for the IMM group at R1 (distal epiphysis), R2 (mid-shaft: distal) R3 (mid-shaft: proximal) and R4 (proximal epiphysis) was significantly higher than those for the normal group, which indicates an immobilization elicited increase in porosity in these areas. The X-ray transmission intensity of ZOL, ALF and PRO groups were significantly lower as compared with IMM group at R1, R2, R3 and R4. The results of the bone porosity analysis had showed no significant difference between all pharmacological treatment or the non-immobilized group.

*Non-immobilized (left) leg*

There were no significant differences between groups concerning bone porosity values (data not shown).
Fig. 3. Effect of ZOL, ALF and PRO on femoral porosity. (a) R1: distal femoral epiphysis, (b) R2: distal femoral shaft, (c) R3: proximal femoral shaft, (d) R4: proximal femoral epiphysis. Data are expressed as the mean ± S.D (n=6), evaluated by one-way ANOVA followed by Tukey’s multiple comparison test.*p < 0.05; **p < 0.01; ***p < 0.001, compared to IMM control group.
Effect of different treatments on mechanical properties of the femoral mid-shaft

Immobilized (right) leg

Fig. 4 (a-e) shows the peak load, ultimate stiffness, fracture toughness, ultimate strength and Young’s modulus in the femoral mid-shaft, respectively. Three-point bending tests of the right femur indicated that immobilization caused significant reductions in the strength parameters including peak load, ultimate stiffness, fracture toughness, ultimate strength and Young’s modulus compared with those in normal or non-immobilized group ($p < 0.001$). Peak load of the femur in ZOL, ALF and PRO groups was significantly higher than in the IMM group ($p < 0.05$, $p < 0.05$ and $p < 0.01$, respectively). Moreover, the ultimate stiffness of the femur in ZOL, ALF and PRO groups was significantly higher than in the IMM group ($p < 0.01$, $p < 0.05$ and $p < 0.01$, respectively). Also the fracture toughness of the femur in the ZOL, ALF and PRO groups was significantly higher than in the IMM group ($p < 0.05$, $p < 0.05$ and $p < 0.01$, respectively). Furthermore, in ZOL, ALF and PRO groups, the ultimate strength of the femur was significantly higher than in the IMM group ($p < 0.05$, $p < 0.05$ and $p < 0.01$, respectively). The Young’s modulus of the ZOL, ALF and PRO groups was significantly increased when compared with the IMM group ($p < 0.01$). In contrast, there were no significant differences among each of the treatment groups with respect to peak load, ultimate stiffness, toughness, ultimate strength and Young’s modulus of the femoral mid-shaft.

Non-immobilized (left) leg

There were no significant differences between peak load, ultimate stiffness, fracture toughness, ultimate strength and Young’s modulus from IMM control animals, normal control animals, or drug-treated animals (data not shown).
(a) PEAK LOAD (N)

(b) ULTIMATE STIFFNESS (N/mm)

(c) TOUGHNESS (mJ)

(d) ULTIMATE STRENGTH (MPa)
Fig. 4. Effects of ZOL, ALF, and PRO on the mechanical strength of the femoral mid-shaft. The femoral mid-shaft was subjected to three-point bending to failure, which provided data on peak load (a), ultimate stiffness (b), toughness (c), ultimate strength (d), and Young’s modulus (e). Data are expressed as the mean ± S.D (n=6), evaluated by one-way ANOVA followed by Tukey’s multiple comparison test.*p < 0.05; **p < 0.01; ***p < 0.001, compared to IMM group.

Scanning Electron Microscopy

At the end of the treatment, compared with the normal group, the cortical structure revealed a higher pore diameter (+37.67%, non-significant) and a higher pore number (+ 41.62 %, p < 0.01) in the IMM group, which yielded an overall higher cortical porosity +78% (p < 0.001). Cortical porosity tended to be lower in ZOL (-70.81%), ALF (-76.95%) and PRO (-79.86%) when compared with IMM group (p < 0.001). Pore number was significantly lower in all treatment groups compared with IMM group. These results were confirmed by those obtained from the
scanning electron micrographs (Fig. 5). Treatment with ZOL, ALF and PRO improved the microarchitecture of bones.
Fig. 5. Scanning electron micrographs taken from femoral proximal-diaphysis. Bone pore sites are shown by arrows. Pore geometry image represents the levels of pore diameter.

Comparison between non-immobilized (Left) leg and immobilized (Right) leg with in a same group

The bone porosity and mechanical properties of the left and right legs are plotted as “split-bar” diagrams in fig. 6 & 7, respectively. An asterisk indicates that there was a significant difference between the left and right leg within the same group (Wilcoxon’s signed-rank test). At R1, R2, R3 and R4 regions, the X-ray transmission intensity for the immobilized side (right) seemed significantly higher than those from the non-immobilized side (left) in the IMM group ($p < 0.01$). In contrast, treatment with the standard therapy or PRO showed full protection against disuse osteoporosis at R1, R2, R3 and R4 regions, as indicated by X-ray transmission intensity values (Fig. 6).

At the femoral mid-diaphysis (three-point bending test), the effect of immobilization was very pronounced in IMM group; that is, the immobilized side (right) had significantly lower values of
strength parameters including peak load, ultimate stiffness, fracture toughness, ultimate strength and Young’s modulus than the non-immobilized side (left) ($p < 0.01$). Treatment with all therapeutic interventions showed full protection against immobilization (Fig. 7).

![Graphs](image-url)

**Fig. 6.** Femoral porosity for the non-immobilized (left bar) and the immobilized (right bar) side with in the same group. Asterisk denotes significant difference between the non-immobilized side and the immobilized side (mean ± SD).
Fig. 7. Mechanical properties for the non-immobilized (left bar) and the immobilized (right bar) side with in the same group. The femoral mid-shaft was subjected to three-point bending to failure, which provided data on peak load (a), ultimate stiffness (b), toughness (c), ultimate strength (d), and Young’s modulus (e). Asterisk denotes significant difference between the non-immobilized side and the immobilized side (mean ± SD).

Discussion

A number of animal models for studying the influence of disuse on bone have been proposed [20]. Immobilization of the hind-limb of rats has been repeatedly used as an animal model to investigate disuse osteoporosis. In laboratory animals, localized disuse of a single extremity can be obtained by different ways: denervation, bone resection, tenotomy, devascularization, or arthrodesis. However, it has been shown that bone loss results from cumulative effects of the disuse and the regional acceleratory phenomenon caused by the surgical trauma. Non-surgical
A method like transient muscle paralysis caused by botulinum toxin A has been found to provoke a rapid bone loss with the cessation of active muscle contraction [21]. Minute quantity of botulinum toxin may spread to adjacent tissues or enter the circulatory system. Due to this diffusion, it can produce regional or systemic side effects [22]. Other non-surgical methods such as immobilization by casting, bandaging or tail suspension are preferred because they do not expose to surgical trauma. The need for special cages, difficulties in obtaining a correct and reliable bandaging, necessity to regularly reapply the tape bandaging (one or two times per week to maintain immobilization), the necessarily redistribution of bone strains due to non-physiological positions, and the muscle compression created in these models limit their interest [20, 23, 24]. Studies conducted in our laboratory showed that the immobilization made of cast are too heavy (50-60 % of the rat weight), so there was difficulty in handling and restraint of the rat during the period of immobilization. Despite the recognized effectiveness of casts in inducing hind-limb immobilization, they must be adjusted to each animal for optimal results, which is very time-consuming. Moreover, a thin layer of padding and a layer of fiberglass must be placed beneath the cast for preventing dermatitis and preventing the animal from chewing through the cast, respectively. Furthermore, the rat must be monitored on a daily basis for fecal clearance, chewed plaster, abrasions, venous occlusion, and problems with ambulation, which may require the readjustment and/or reinforcement of the cast. Other kind of difficulties with this technique of immobilization involves a series of complicating factors, i.e, difficulty in observing the rat skin and in some cases rat skin showed skin ulceration or swelling probably due to the retention of urine by the cast. In some cases, rats showed a marked loss of weight due to fixed immobilization and development of edema in the distal extremity of the immobilized limb. Moreover, some rats were able to slip out of the immobilization. Since cast immobilization is a
major experimental challenge, the development of a convenient, easy-to-perform immobilization procedure will significantly facilitate this important area of research.

After several tests trying to find an alternative method for immobilization to avoid the problems caused by a cast, we developed a new model which is described in this research paper. The new immobilization model adopted in the present study has shown that, in rats, 10 weeks of right hind-limb immobilization caused a very pronounced loss of bone. In the data obtained in preliminary studies carried out in our laboratory, the smaller weight of the frame work compared with a plaster cast, kept the difficulty in movement and locomotion to a minimum, with a consequent minimal body weight loss throughout the period of immobilization. Moreover, no skin ulceration or foot swelling was found in the animals when the immobilization was removed.

Body weight in the normal group was greater than in the IMM group. This indicates that the decrease in body weight of IMM group was caused by right hindlimb immobilization. Earlier studies by others have shown a similar decrease in body weight after immobilization [2, 25].

Kondo et al indicated that sympathetic tones regulate immobilization-induced enhancement in bone resorption. This was evidenced by the observations that inhibition of sympathetic tone by propranolol, suppressed immobilization-induced bone resorption, and this leads to suppression in bone loss [17]. As expected, PRO administered for 10 weeks reduced the osteopenia induced by immobilization, as demonstrated by the improvements in bone properties in immobilized long bones. Interestingly, immobilization of the right leg also induced a small decrease in bone properties of the non-immobilized contralateral leg, most likely because the immobilized animals
did not use that leg as much as the control animals. Moreover, in the present study, osteoporosis induced by immobilization did not result from alteration of skeletal growth because body growth and longitudinal femur length in the immobilized limb were not altered.

Microarchitectural observation of fractured surfaces of femur (proximal-diaphysis) was performed with SEM. Results of SEM analysis showed poor microarchitecture of bone (femur) with increase in cortical porosity in IMM control animals as against normal control. Rats treated with PRO had lower cortical porosity, pore number, and higher space between pores compared with the IMM group. The current data correlate with findings from a previous study conducted by Bonnet et al [19], demonstrating effects of PRO on cortical bone properties.

In this study, prominent increase in porosity was observed at R1 (femoral distal epiphysis), R2 (mid-shaft: distal), R3 (femoral mid-shaft: proximal), and R4 (femoral proximal epiphysis), after immobilization. The increase in the bone porosity at R1-R4 regions of rat femoral bone, due to unloading was suppressed by treatment with PRO. Similarly, in ZOL and ALF groups the protective effect on femoral porosity was observed at R1, R2, R3, and R4 regions. The results of bone porosity found by X-ray imaging were consistent with those on microarchitectural observation of fractured surfaces of femur (proximal-diaphysis) by SEM.

Unloading induces rapid bone loss and increases fracture risk significantly, especially in elderly bedridden patients. In this experiment, the mechanical properties of the rat femoral bone decreases in IMM group when compared to normal group, suggesting an increase in the fragility of the cortical bone of IMM rats. PRO was capable of maintaining bone strength at a normal
level (not significantly different from the non-immobilized leg). Similarly, ZOL and PRO maintained bone strength at the distal femoral metaphysis. It should be noted, though, that there were no significant differences in various bone parameter values in a direct comparison between PRO, ZOL and ALF. The current data correlate with findings from our previous study on PRO, demonstrating effects of PRO on the mechanical properties of ovariectomized rat bone [9].

The present study has also clearly demonstrated that the two standard drugs, ZOL and ALF, each effectively protected against immobilization-induced bone loss. ZOL and ALF were capable of maintaining bone porosity, cortical microarchitecture and bone strength at a normal level (not significantly different from the non-immobilized leg). ZOL therapy, which is the popular regime for osteoporosis, is associated with increased risk of osteonecrosis of jaw, atrial fibrillation, atypical fracture and renal toxicity. Similarly, treatment with ALF is associated with higher risk of hypercalcemia and hypercalciuria [6]. Studies have demonstrated that that heart hemodynamic functions were preserved under 0.1 mg/kg/day dose of PRO treatment. Cardiac output and other echocardiographic assessments were unaffected by low dose of PRO, the same dose that prevented bone loss [10, 13]. If this is the case, PRO, with little known toxicity, would make an excellent candidate for the therapeutic prevention and treatment of disuse osteoporosis.

Conclusion

In summary, the immobilization framework proposed in this study was effective in producing long-term disuse in the hind-limbs of rats and is a good alternative to the traditional methods of immobilization. Present study tentatively suggests that PRO is effective treatment of immobilization osteoporosis. The findings are consistent with the effects of PRO on estrogen
deficiency bone loss and extend our knowledge regarding the effects of this therapy in immobilization-induced bone loss. The present study indicates that PRO may play an important role in the clinical management of osteoporosis and similar disorders.

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Authors’ Statements
Competing Interests
The authors declare no conflict of interest.

Animal Rights
The institutional and (inter)national guide for the care and use of laboratory animals was followed. See the 'materials and methods' part for details.
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