Combination therapy with low-dose teriparatide and zoledronate contributes to fracture healing on rat femoral fracture model

Yuta Tsubouchi, Shinichi Ikeda, Masashi Kataoka and Hiroshi Tsumura

Abstract

Background: Delay in fracture healing or non-union can be devastating complication. Recent studies have reported that teriparatide (TP) demonstrated effectively on callus formation and mechanical strength and zoledronate (ZA) increased the callus size and resistance at the fracture site in rat fracture model. In this study, the effects of combination therapy with low dose TP and ZA on fracture healing was evaluated.

Methods: From 1 week post-operation, TP (5 times a week administration) and ZA (0.1 mg/kg single administration) were administered by dividing the rats into the following five groups: TP 1 μg group (T(1): TP 1 μg/kg), ZA group (ZA:0.1 mg/kg), TP1 μg+ZA group (T(1)+ZA: TP 1 μg/kg+ZA), TP 10 μg+ZA group (T(10)+ZA: TP 10 μg/kg + ZA), and control group (C: administered saline). Rt femurs were excised 7 weeks after the surgery; bone fusions were evaluated with soft X-ray images on a 4-point scale. And the histopathological examination was performed in demineralized and non-demineralized specimens. Furthermore, the Radiographic Union Scale was conducted in all specimens.

Results: About the bone fusions rates, C, T(1), ZA, T(1)+ZA, and T(10)+ZA groups demonstrated 20.0%, 55.6%, 70.0%, 70.0%, and 80.0%, respectively, and with 4-point scale, each group was 0.50, 1.56, 2.00, 2.60, and 2.80 points, respectively. The callus volume was significantly increased to 16.66 mm$^2$ and 17.75 mm$^2$ in the T(1)+ZA and T(10)+ZA groups, respectively, while 10.65 mm$^2$ ($p < 0.05$) in the C group. Furthermore, the callus area in the T(10)+ZA group was also observed to have significantly increased to 78.78%, compared with 54.63% and 44.11% in the C and T(1)+ZA groups, respectively ($p < 0.01$). Histopathologically, cartilage tissue and immature callus formation were observed at the bone junction in the C group; however, the osseous bridge formation of mature callus was observed in the ZA, T(1)+ZA, and T(10)+ZA groups.

Conclusion: It is suggested that administration of low dose TP and ZA in combination may lead to the treatment of delayed union of fracture. We hope the combination treatment may become one of new therapeutic strategy.

Keywords: Fracture healing, Teriparatide, Zoledronate
Background
Although improvements in surgical methods for fractures of the femoral shaft have resulted in delayed fracture healing and nonunion in up to 5–10% of patients, subsequent treatment upon the occurrence of nonunion is challenging. Achieving early bone union and returning patients to society under such unfavorable conditions are important issues to address [1].

Teriparatide (TP), which was used in the present study, affects osseous tissue, and its effect varies according to the administration method. While a catabolic effect on the bone is observed during continuous administration, an anabolic effect occurs when it is intermittently administered. Since 2010, TP has been widely used as a therapeutic drug for osteoporosis in Japan; however, it is expected to be clinically applied in the future as a therapeutic agent that promotes fracture healing because of its intense osteogenesis-promoting effect. Coppola et al. reported that TP administration was really effective for the management of non-unions in four cases; however, the efficacy of the TP in delayed or non-unions was not clear [2]. In a rat femoral fracture model, Andreassen et al. administered a post-fracture subcutaneous injection of parathyroid hormone 1–34 (PTH 1–34) for consecutive days at two doses (60 and 200 μg/kg) and reported greater callus formation and strong callus in the group administered 200 μg/kg/PTH [3]. However, the continuous administration of 60 or 200 μg/kg PTH is unrealistic and poses pharmacological problems. Considering the administration to humans in clinical practice, examinations should be conducted using a lower dosage. Several studies have investigated the administration of low-dose rhPTH (1–34; 10 μg/kg) in a rat femoral fracture model, in which treatment increased callus bone mineral content (BMC) and bone density, as well as dynamic strength and was thus found to be effective [4, 5].

In contrast, bisphosphonate (BP) has a bone resorption inhibitory action and has been demonstrated to effectively improve bone mineral density (BMD), normalize elevated levels of serum bone metabolism markers, and lower the risk of bone fractures [6, 7]. Furthermore, several reports have studied the effects of BP on the process of bone fracture healing. Most of these reports have demonstrated effectiveness in increasing callus volume (CV) and dynamic strength [8, 9]. In a study conducted to investigate the effects of 0.1-mg/kg zoledronate (ZA) administered in an ovariectomized rat femoral fracture model, it was reported that ZA increased callus strength, bone mass:volume ratio, trabecular width, and trabecular connectivity density during the process of bone fracture healing [10].

Purpose
Although there are not many studies on the effect of combined TP and ZA therapy on the process of bone fracture healing, the effectiveness of this treatment has been reported; it increases bone mass:volume ratio of the callus as well as bone mass and width [11, 12]. These studies were conducted with the administration of high-dose PTH, which cannot be considered a realistic dose in clinical practice. Therefore, the present study aimed to examine the effect of combined low-dose TP and ZA for bone fracture healing in a refractory rat fracture model using X-ray evaluation and histopathological evaluations of demineralized and non-demineralized specimens.

Methods
Study groups
A total of 50 male Sprague–Dawley rats (12 weeks old; CLEA Japan, Inc., Tokyo, Japan) were divided into five groups. From 1 week post-operation, TP (Forteo, Lilly Japan, Tokyo) (5 times a week administration) and ZA (Novartis Pharma KK Tokyo Japan) (0.1 mg/kg single administration) were administered by dividing the rats into the following 5 groups: TP 1 μg group (T1): TP 1 μg/kg, ZA group (ZA:0.1 mg/kg), TP1 μg+ZA group (T1+ZA:TP 1 μg/kg+ZA), TP10 μg+ZA group (T10+ZA: TP 10 μg/kg+ZA), and control group (C: administered saline).

Surgical technique used to construct the femoral fracture model
Approval was obtained from the Oita University animal research committee prior to animal experimentation (Oita University institutional Animal Ethics Committee no.1624003). The Sprague–Dawley rats were anesthetized by an intraperitoneal injection containing 0.3–0.4 ml of 0.15 mg/kg medetomidine + 2 mg/kg midazolam + 2.5 mg/kg butorphanol. The right hind limb was prepared for the operation under standard sterile conditions. With the rat in the lateral position, the right femur was located using the posterolateral approach. The periosteum of the femur was circumferentially incised and elevated and stripped. Then, the femur at osteotomy site was exposed. A transverse osteotomy was performed at the mid-shaft of the femoral bone, the fracture fragments were contacted and stabilized, and the intramedullary was then fixed using a stainless steel wire (diameter, 1.6 mm). The wire was cut on the surface of the intercondylar groove to avoid knee joint motion restriction. The material was applied and wrapped circumferentially around the fracture site. The fascial and skin incisions were closed with a 3–0 nylon suture.

Immediately following surgery and on subsequent days, the rodents received analgesics (buprenorphine subcutaneously and paracetamol). The rodents were housed in separate cages and fed food and water ad libitum, and their conditions were monitored daily. The rats were humanly euthanized 6 weeks after the operation,
and the operated right femoral bones were explanted and separated from the stainless steel wire before analysis. We used the left femoral bones, which had not been operated upon, as controls in the biomechanical analysis.

Radiographic analysis
The explanted femoral bones obtained at the 7-week time point were photographed using a Softex X-ray apparatus (Softex CSM-2; Softex, Tokyo, Japan) employing HS Fuji Softex film (Fuji Film, Tokyo, Japan) at 45 cm with 30 kV and 15 mA for 20 s. The fusion was quantified using anteroposterior (A-P) and lateral radiographs. Three blinded independent observers scored the bone formation in each rat using a 4-point scale. Fracture union was judged by visual assessment of the mineralized callus bridging the fracture line in the A-P radiographs (right side: 1 point; left side: 1 point) and lateral radiographs (anterior side: 1 point; posterior side: 1 point). The bone fusion was considered to be more than 2 points with soft X-ray images on a 4-point scale. Furthermore, the Radiographic Union Scale for Tibial fractures in Tibial Fracture (RUST) was conducted in all specimens [13]. The RUST score is based on the presence or absence of callus and of a visible fracture line at the total of four cortices visible on the anteroposterior and lateral radiographs. Its 4-point minimum corresponds to a fracture that is deemed not healed, whereas its 12-point maximum corresponds to a fracture that is deemed healed with all cortices bridged with callus without a fracture line.

Histological analysis
Ten specimens of one group were divided into two kinds of staining. For the HE staining, 4 of 10 specimens were performed, and for the non-decalcified and toluidine blue staining, 6 of 10 specimens were performed in a random manner. After extraction, the femoral bones were dissected, and the specimens were fixed in 70% ethanol. The specimens were then decalcified using a 10% decalcifying solution of HCl (Cal-Ex) (Fisher Scientific, Fairlawn, NJ), washed with running tap water, and transferred to 75% ethanol. Serial sagittal sections (5 μm) were carefully cut from the paraffin blocks using a microtome (LS-113; DAIWA-KOKI, Saitama, Japan) at the level of the femoral fracture. The sections were stained with hematoxylin and eosin (HE) and evaluated qualitatively under a light microscope (n = 4 per groups).

Furthermore, non-decalcified specimens were obtained from glycolmenthacrylate resin-embedded slices, and the calcification state and cells in the fracture area were observed (n = 6 per groups). After staining with toluidine blue, bone morphometry was performed to up to 0.43 mm proximal and distal to the bone fracture area (Histometry RT CAMERA; System Supply, Tokyo, Japan). Measurement items included callus volume (CV), callus area (CA), and cartilage tissue area (Ca.A).

Statistical methods
One-way ANOVA with Bonferroni post hoc test was used for 4-point test and RUST and the analysis of morphometric results of the treatment groups. All analyses were performed using Statistical Package for the Social Sciences (SPSS V22.0; SPSS, Chicago, IL). Statistical significance was set at < 0.05.

Results

Radiographic analysis
In the C and T(1) group, the fracture line could be clearly identified, and immature callus formation was observed. In particular, callus continuity was not achieved in the C group. Conversely, in the ZA, T(1)+ZA, and T(10)+ZA groups, the fracture line was unclear, the callus was thick, and continuity was confirmed (Fig. 1). On the 4-point scale, the C group scored 0.50 ± 0.85 points, the T(1) group scored 1.56 ± 1.67 points, the ZA group scored 2.00 ± 1.41 points, the T(1)+ZA group scored 2.60 ± 1.90 points, and the T(10)+ZA group scored 2.80 ± 1.69 points, indicating significantly more progress in bone union in the T(1)+ZA and T(10)+ZA groups than in the C group (Table 1).

The RUST score results, including the evaluation of cortical bone continuity and callus formation, were comparable among the groups, with the C group scoring 5.00 ± 0.82 points, the T(1) group scoring 6.67 ± 2.24 points, the ZA group scoring 7.90 ± 3.10 points, the T(1)+ZA group scoring 9.20 ± 3.65 points, and the T(10)+ZA group scoring 10.2 ± 2.57 points, indicating significantly more progress in bone union in the T(1)+ZA and T(10)+ZA groups than in the C group (Fig. 2).

Bone histomorphometrical analysis
In the C and T(1) group, verifying the fracture site by HE staining identified the fracture and confirmed extensive chondrogenesis. Furthermore, while several specimens demonstrated thin immature callus formation, callus continuity could not be confirmed in many specimens. Although a few chondrocytes were identified in the T(1)+ZA group, they were enlarged. Compared to the C group, the fracture line was unclear. Furthermore, in the T(10)+ZA group, chondrocytes could not be identified, but mature callus and advanced bone union were confirmed (Fig. 3).

Staining with toluidine blue revealed cartilage tissue and immature callus formation along with the fracture

...and the operated right femoral bones were explanted and separated from the stainless steel wire before analysis. We used the left femoral bones, which had not been operated upon, as controls in the biomechanical analysis.

Radiographic analysis
The explanted femoral bones obtained at the 7-week time point were photographed using a Softex X-ray apparatus (Softex CSM-2; Softex, Tokyo, Japan) employing HS Fuji Softex film (Fuji Film, Tokyo, Japan) at 45 cm with 30 kV and 15 mA for 20 s. The fusion was quantified using anteroposterior (A-P) and lateral radiographs. Three blinded independent observers scored the bone formation in each rat using a 4-point scale. Fracture union was judged by visual assessment of the mineralized callus bridging the fracture line in the A-P radiographs (right side: 1 point; left side: 1 point) and lateral radiographs (anterior side: 1 point; posterior side: 1 point). The bone fusion was considered to be more than 2 points with soft X-ray images on a 4-point scale. Furthermore, the Radiographic Union Scale for Tibial fractures in Tibial Fracture (RUST) was conducted in all specimens [13]. The RUST score is based on the presence or absence of callus and of a visible fracture line at the total of four cortices visible on the anteroposterior and lateral radiographs. Its 4-point minimum corresponds to a fracture that is deemed not healed, whereas its 12-point maximum corresponds to a fracture that is deemed healed with all cortices bridged with callus without a fracture line.

Histological analysis
Ten specimens of one group were divided into two kinds of staining. For the HE staining, 4 of 10 specimens were performed, and for the non-decalcified and toluidine blue staining, 6 of 10 specimens were performed in a random manner. After extraction, the femoral bones were dissected, and the specimens were fixed in 70% ethanol. The specimens were then decalcified using a standard 10% decalcifying solution of HCl (Cal-Ex) (Fischer Scientific, Fairlawn, NJ), washed with running tap water, and transferred to 75% ethanol. Serial sagittal sections (5 μm) were carefully cut from the paraffin blocks using a microtome (LS-113; DAIWA-KOKI, Saitama, Japan) at the level of the femoral fracture. The sections were stained with hematoxylin and eosin (HE) and evaluated qualitatively under a light microscope (n = 4 per groups).

Furthermore, non-decalcified specimens were obtained from glycolmenthacrylate resin-embedded slices, and the calcification state and cells in the fracture area were observed (n = 6 per groups). After staining with toluidine blue, bone morphometry was performed to up to 0.43 mm proximal and distal to the bone fracture area (Histometry RT CAMERA; System Supply, Tokyo, Japan). Measurement items included callus volume (CV), callus area (CA), and cartilage tissue area (Ca.A).

Statistical methods
One-way ANOVA with Bonferroni post hoc test was used for 4-point test and RUST and the analysis of morphometric results of the treatment groups. All analyses were performed using Statistical Package for the Social Sciences (SPSS V22.0; SPSS, Chicago, IL). Statistical significance was set at < 0.05.

Results

Radiographic analysis
In the C and T(1) group, the fracture line could be clearly identified, and immature callus formation was observed. In particular, callus continuity was not achieved in the C group. Conversely, in the ZA, T(1)+ZA, and T(10)+ZA groups, the fracture line was unclear, the callus was thick, and continuity was confirmed (Fig. 1). On the 4-point scale, the C group scored 0.50 ± 0.85 points, the T(1) group scored 1.56 ± 1.67 points, the ZA group scored 2.00 ± 1.41 points, the T(1)+ZA group scored 2.60 ± 1.90 points, and the T(10)+ZA group scored 2.80 ± 1.69 points, indicating significantly more progress in bone union in the T(1)+ZA and T(10)+ZA groups than in the C group (Table 1).

The RUST score results, including the evaluation of cortical bone continuity and callus formation, were comparable among the groups, with the C group scoring 5.00 ± 0.82 points, the T(1) group scoring 6.67 ± 2.24 points, the ZA group scoring 7.90 ± 3.10 points, the T(1)+ZA group scoring 9.20 ± 3.65 points, and the T(10)+ZA group scoring 10.2 ± 2.57 points, indicating significantly more progress in bone union in the T(1)+ZA and T(10)+ZA groups than in the C group (Fig. 2).

Bone histomorphometrical analysis
In the C and T(1) group, verifying the fracture site by HE staining identified the fracture and confirmed extensive chondrogenesis. Furthermore, while several specimens demonstrated thin immature callus formation, callus continuity could not be confirmed in many specimens. Although a few chondrocytes were identified in the T(1)+ZA group, they were enlarged. Compared to the C group, the fracture line was unclear. Furthermore, in the T(10)+ZA group, chondrocytes could not be identified, but mature callus and advanced bone union were confirmed (Fig. 3).

Staining with toluidine blue revealed cartilage tissue and immature callus formation along with the fracture
line in the C and T(1) groups, whereas osseous bridge formation of the mature callus was observed in the ZA, T(1)+ZA, and T(10)+ZA groups. Furthermore, bone union was seen resulting from endochondral ossification in the ZA, T(1)+ZA, and T(10)+ZA groups, and mature bone tissue was identified in the T(10)+ZA group (Fig. 1). Bone histomorphometry revealed CV was 10.65 ± 2.47 mm² in the C group, 8.01 ± 1.46 mm² in the T(1) group, and 12.79 ± 1.28 mm² in the ZA group, but was significantly higher at 16.66 ± 2.35 mm² in the T(1)+ZA group and 17.75 ± 2.85 mm² in the T(10)+ZA group (P < 0.01). Similarly, CA was 54.63% ± 5.31% in the C group, 44.11% ± 4.13% in the T(1) group, 62.84% ± 2.75% in the T(1)+ZA group, and 62.57% ± 2.75% in the T(10)+ZA group, but it could not be confirmed in the ZA and T(10)+ZA group (P < 0.05) (Fig. 4a, b, c).

Discussion

Einhorn et al. reported the molecular mechanism of fracture healing in rodents from a molecular biological perspective. Evaluation of femoral shaft fracture was performed with Softex X-ray apparatus. Fracture union was judged by visual assessment of the mineralized callus bridging the fracture line in the A-P radiographs (right side: 1 point; left side: 1 point) and lateral radiographs (anterior side: 1 point; posterior side: 1 point). The bone fusion was considered to be more than 2 points with soft X-ray images on a 4-point scale. About the bone fusion rates, C, T(1), ZA, T(1)+ZA group, and T(10)+ZA group demonstrated 20.0%, 55.6%, 70.0%, 70.0%, and 80.0% respectively. With the 4-point scale analysis, T(10)+ZA and T(1)+ZA groups were significantly superior to C group in bone union.

Table 1 Results of bone radiographic statistical analysis of 4-point scale and bone fusion rate

|                | Control (n = 10) | T(1) (n = 10) | ZA (n = 10) | T(1)+ZA (n = 10) | T(10)+ZA (n = 10) | ANOVA  |
|----------------|-----------------|---------------|-------------|------------------|-------------------|--------|
| Fusion rate (%)| 20.0            | 55.6          | 70.0        | 70.0             | 80.0              |        |
| 4-point scale  | 0.50 ± 0.85     | 1.56 ± 1.67   | 2.00 ± 1.41 | 2.60 ± 1.90      | 2.80 ± 1.69       | 0.014  |

All values are mean ± standard deviation

Evaluation of femoral shaft fracture was performed with Softex X-ray apparatus. Fracture union was judged by visual assessment of the mineralized callus bridging the fracture line in the A-P radiographs (right side: 1 point; left side: 1 point) and lateral radiographs (anterior side: 1 point; posterior side: 1 point). The bone fusion was considered to be more than 2 points with soft X-ray images on a 4-point scale. About the bone fusion rates, C, T(1), ZA, T(1)+ZA group, and T(10)+ZA group demonstrated 20.0%, 55.6%, 70.0%, 70.0%, and 80.0% respectively. With the 4-point scale analysis, T(10)+ZA and T(1)+ZA groups were significantly superior to C group in bone union.

T(1) teriparatide 1 µg group, ZA zoledronate group, T(1)+ZA teriparatide 1 µg + zoledronate group, T(10)+ZA teriparatide 10 µg + zoledronate group

*p < 0.05 vs. control group, by post hoc test
TGF-β and IL-6 were derived from inflammatory cells, and T(1)+ZA and T(10)+ZA groups than in the C group, indicating significantly more progress in bone union in the T(1)+ZA and T(10)+ZA groups than in the C group. We had performed several times of experiments of femoral fracture model, and the average rate of bone union was almost 10–20% of the cases in stripping of peristeum. Cartilage formation is taking place in the cambium layer of the peristeum, and chondrocyte precursor cells also reside in this location. Accordingly, bone union must rely on endochondral ossification. To date, studies have illustrated that TP promotes endochondral ossification from cartilage formation occurring in the fracture gap and promotes membranous ossification near the fracture site. Furthermore, TP promotes callus remodeling and reduces the bridging callus, simultaneously increases the degree of calcification, improves the dynamic quality of callus, and promotes a normal bone fracture healing process.

Nakazawa et al. injected 10 μg/kg of PTH in a rat closed femoral fracture model and analyzed the promotion of bone fracture healing at the molecular level [5]. Compared to the control group, the group administered with PTH radiologically demonstrated remarkable callus formation from the early stage to the late stage in the healing process and exhibited a significantly higher BMC, bone density, and dynamic strength at 4 and 6 weeks after fracture. Several studies have examined bone fracture healing in animals treated with BP. The results of these studies revealed that BP accumulated in the acute fracture site predominantly during the repair phase of bone healing. Furthermore, it was determined that BP not only helped increase callus and bone mass but also increased BMC. Conversely, BP did not affect the start of callus formation.

Regarding the effect of BP on endochondral ossification, it has been suggested that the size and strength of the callus can be controlled by regulating bone remodeling during the healing process. Consequently, the timing and regimen of BP therapy can have a significant impact on the formation and strength of the callus. In our preliminary experiment, we established that 1 week after fracture is the optimal timing of a single intraperitoneal injection of ZA for the rate of bone union, bone formation, and strength testing [15]. Furthermore, Murphy et al. reported that PTH (1–34) increased bone turnover as well as BP turnover [16]. Treatment of combined PTH and BP may be increased effect of BP.

Histopathological findings of the species revealed that ZA treatment not only increases callus formation but also improves trabecular microarchitecture. ZA given systemically as a single dose at the optimal time could reduce catabolic osteoclastic resorption caused by TP.
However, there were no evidence of ZA-induced suppression of anabolic bone formation caused by TP. The fusion rates in rat femurs administered both ZA and TP were higher than those of rats administered ZA only 6 weeks after surgery.

The present study demonstrated that the synergistic effect of low-dose TP and ZA promoted callus formation, increased callus mass, and helped to promote bone union in a rat refractory fracture model. The synergistic effect of the combination of TP with the anabolic effect and ZA with the anti-catabolic effect was also demonstrated. Because ZA promotes the apoptosis of osteoclasts, remodeling is delayed, which, in turn, can impede the change from fibrous bone to lamellated bone. Hence, detailed

---

**Fig. 3** Bone histological imaging (demineralized specimens: HE staining) (×1) (×40). In the C (a) and T(1)+ZA groups (b), verifying the fracture site by HE staining identified the fracture and confirmed extensive chondrogenesis. Furthermore, while several specimens demonstrated thin immature callus formation, callus continuity could not be confirmed in many specimens. And cartilaginous bone metaplasia was not observed. Although a few chondrocytes, suggesting bone metaplasia, were identified in the T(1)+ZA group (b), they were enlarged. Compared to the C group, the fracture line was unclear. Furthermore, in the T(10)+ZA group (c), chondrocytes could not be identified, but mature callus and advanced bone union were confirmed (c).

**Fig. 4 a, b, c** Results of bone histomorphometry. a Callus volume (CV). b Callus area (CA). c Cartilage tissue area (Ca.A). Bone histomorphometry revealed callus volume (CV): C 10.65 ± 2.47 mm², T(1) 8.01 ± 1.46 mm², and ZA group 12.79 ± 1.28 mm², but was significantly higher at 16.66 ± 2.35 mm² in the T(1)+ZA group and 17.75 ± 2.85 mm² in the T(10)+ZA group (P < 0.01). Similarly, CA was 54.63% ± 5.31% in the C group, 44.11% ± 4.13% in the T(1) group, 62.84% ± 5.89% in the ZA group, and 62.57% ± 2.75% in the T(1)+ZA group, but was significantly higher at 78.78% ± 0.85% in the T(10)+ZA group (P < 0.001). However, Ca.A was 0.74 ± 0.51 mm² in the C group, 0.06 ± 0.11 mm² in the T(1) group, 0.01 ± 0.02 mm² in the T(1)+ZA group, but it could not be confirmed in the ZA and T(10)+ZA group (p < 0.05).
investigations of the bone structure through long-term observations are required in the future.

**Conclusion**

The present experiment shows that administration of low-dose TP and ZA in combination is able to enhance callus formation, increase callus volume, and help to promote bone union in a rat refractory fracture model. We hope the combination treatment may become one of new therapeutic strategy in delay in fracture healing or non-union of patients.

**Abbreviations**

BMC: Bone mineral content; BMD: Bone mineral density; BP: Bisphosphonate; CA: Callus area; Ca.A: Cartilage tissue area; CV: Callus volume; PTH: Parathyroid hormone; RUST: The Radiographic Union Scale in Tibial Fracture; TP: Teriparatide; ZA: Zoledronate

**Acknowledgements**

We are grateful to Miss Kaori Abe for excellent technical assistance.

**Funding**

This work was supported by JSPS KAKENHI Grant Number JP16K12939.

**Availability of data and materials**

All data generated or analyzed during this study are included in this published article.

**Authors' contributions**

YT, SI, and MK carried out the operation. HT conceived of the study design. YT, SI, MK, and HT interpreted the data and drafted the manuscript. All authors read and approved the final manuscript.

**Ethics approval**

This study was conducted in accordance with the Declaration of Helsinki. Approval was obtained from the Oita University animal research committee prior to animal experimentation (Oita University institutional Animal Ethics Committee no.1624003).

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

**References**

1. Buza JA 3rd, Einhorn T. Bone healing in 2016. Clin Cases Miner Bone Metab. 2016;13(2):101–5.
2. Coppola C, Del Buono A, Maffulli N. Teriparatide in fracture non-unions. Trans Med UniSa. 2015;12(8):47–53.
3. Andreasen TT, Ejstrup C, Oxlund H. Intermittent parathyroid hormone (1–34) treatment increases callus formation and mechanical strength of healing rat fractures. J Bone Miner Res. 1999;14(6):960–8.
4. Nakajima A, Shimoyi N, Shimoi K, Shimizu S, Moriya H, Einhorn TA, Yamazaki M. Mechanisms for the enhancement of fracture healing in rats treated with intermittent low-dose human parathyroid hormone (1–34). J Bone Miner Res. 2002;17(11):2038–47.
5. Nakazawa T, Nakajima A, Shimoi K, Moriya H, Einhorn TA, Yamazaki M. Effects of low-dose, intermittent treatment with recombinant human parathyroid hormone (1–34) on chondrogenesis in a model of experimental fracture healing. Bone. 2003;37(1):711–9.
6. Chapurlat RD, Palermo L, Ramsay P, Cummings SR. Risk of fracture among women who lose bone density during treatment with alendronate. The fracture intervention trial. Osteoporos Int. 2005;16(7):842–8.
7. Rebollodo BJ, Unnanuntana A, Lane JM. A comprehensive approach to fragility fractures. J Orthop Trauma. 2011;25(9):566–73.
8. Amanat N, McDonald M, Godfrey C, Bilston L, Little D. Optimal timing of a single dose of zoledronic acid to increase strength in rat fracture repair. J Bone Miner Res. 2007;22(6):867–76.
9. Matos MA, Tannuri U, Guarniero R. The effect of zoledronate during bone healing. J Orthopaed Traumatol. 2010;11(1):7–12.
10. Hao Y, Wang X, Wang L, Lu Y, Mao Z, Ge S, Dai K. Zoledronic acid suppresses callus remodeling but enhances callus strength in an osteoporotic rat model of fracture healing. Bone. 2015;81:702–11.
11. Li YF, Zhou CC, Li JH, Luo E, Zhu SS, Feng G, Hu J. The effects of combined human parathyroid hormone (1-34) and zoledronic acid treatment on fracture healing in osteoporotic rats. Osteoporos Int. 2012;23(4):1463–74.
12. Casanova M, Herelle J, Thomas M, Softley R, Schindeler A, Little D, Schneider P, Müller R. Effect of combined treatment with zoledronic acid and parathyroid hormone on mouse bone callus structure and composition. Bone. 2016;92:70–8.
13. Kooistra BW, Dijkman BG, Busse JW, Sprague S, Schensttisch EH, Bhandari M. The radiographic union scale in tibial fractures: reliability and validity. J Orthop Trauma. 2010;24(Suppl 1):S81–6.
14. Einhorn TA, Gerstenfeld LC. Fracture healing: mechanisms and interventions. Nat Rev Rheumatol. 2015;11(1):45–54.
15. Doi Y, Miyazaki M, Yoshiwa T, Hara K, Kataoka M, Tsumura H. Manipulation of the anabolic and catabolic responses with BMP-2 and zoledronic acid in a rat femoral fracture model. Bone. 2011;49(4):777–82.
16. Murphy CM, Schindeler A, Cantrill LC, Milukcev K, Peacock L, Little DG. PTH (1-34) treatment increases bisphosphonate turnover in fracture repair in rats. J Bone Miner Res. 2015;30(6):1022–9.