Effect of Teriparatide on Healing of Atypical Femoral Fractures: A Systemic Review

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Background: Bisphosphonates (BPs) are the most commonly used anti-osteoporotic drugs, which have been proven to reduce the risk of osteoporotic fractures. However, use of BPs, particularly for long periods of time, is associated with an increased risk of atypical femoral fracture (AFF). Healing of BP-associated AFF is usually delayed because of suppressed bone turnover. Teriparatide (TPTD), a recombinant form of parathyroid hormone (PTH), enhances bone healing in patients with delayed healing or non-union.

Methods: In this study, we summarized and performed a systemic review of the published literature on treatment of AFF using TPTD. Results: Although there is a lack of level 1 studies on the evidence of TPTD in promoting bone union in AFFs, this systemic review of the available literature revealed that TPTD works positively in AFFs, and we put together the evidence that TPTD is a viable treatment option for enhancing fracture healing in AFFs.

Conclusions: While anecdotal evidence of beneficial effects of TPTD on fracture healing offer limited guidance for clinical decision making, a better understanding of the role of TPTD in fracture healing may be elucidated with future prospective trials.

Key Words: Bisphosphonates, Femoral fractures, Parathyroid hormone, Teriparatide

INTRODUCTION

Bisphosphonates (BPs) are the most commonly used anti-osteoporotic drugs, which have been proven to reduce the risk of osteoporotic fracture. BPs are also known to improve quality of life in patients with osteoporosis [1] and reduce mortality in patients who have sustained hip fractures.[2] However long term use of BPs is associated with an increased risk of atypical femoral fracture (AFF).[3,4] AFFs are transverse or oblique stress fractures without comminution occurring in the cortex of the subtrochanteric region or shaft of femur.[5-7]

Anti-resorptive agents including BPs reduce the rate of bone remodeling, therefore slowing the progression of structural loss in bone matrix. Bone matrix then undergoes more complete secondary mineralization rather than being removed and replaced with younger and less mineralized bone matrix,[8] which paradoxically makes bones more brittle because it cannot absorb energy by deformation when loaded. Subsequently, the energy applied to the bone is dissipated by micro-cracking.[9] As seen in prolonged BP administration, microcracks propagate and lengthen with less resistance in homogeneously mineralized bone matrix.[10] In addition, reduced removal of bone microdamage increases occurrence and
propagation of microcracks and compromises the material strength of bone, leading to increased occurrence of stress fractures.[11]

Because of suppressed bone turnover, healing of BP-associated AFF is also usually delayed.[12,13] Teriparatide (TPTD), a recombinant form of parathyroid hormone (PTH) and an anti-osteoporotic agent with potent bone-forming effects,[14,15] enhances bone healing in patients with delayed healing or non-union.[16] A number of studies have demonstrated beneficial effects of TPTD on fracture healing in various animal models, both radiographically and histologically.[17-21] However, there is only limited data for humans currently available for TPTD in fracture healing. [22] When AFF is diagnosed in a patient, the use of BPs as anti-osteoporotic drug is stopped and replaced with TPTD or a selective estrogen receptor modulator. The positive effect of TPTD on fracture healing has prompted the use of TPTP to promote the union of AFF as well.[23,24] Although a number of narrative literature reviews exist elaborating on the role of TPTD in AFF, there is a lack of systematic review of the literature on the subject. In this study, we summarized and performed an in-depth review from published literature on the treatment of AFF using TPTD.

**METHODS**

We performed searches of PubMed, EMBASE, Web of Science, and the Cochrane Database of Systematic Reviews (CDSR) using the search terms “TPTD”, “PTH”, “atypical fracture”, “atypical femoral fracture”, and “healing.” We also screened the references of the searched articles for additional information. Due to the limited amount of literature available on this topic, case reports and case series were also included in the final analysis. Exclusion criteria included articles not relevant to the use of TPTD in AFF, non-English articles, stand-alone abstracts, meeting presentations, commentaries, and review articles.

Searching the aforementioned databases yielded 72 results. Two authors independently read and reviewed the abstracts and full text of the retrieved articles. Eleven articles were deemed appropriate for our study. These included 1 article reporting results from a prospective study, 3 articles from retrospective case series, and 7 case reports. These articles were then analyzed in detail (Fig. 1).

**RESULTS**

Gomberg et al.[25] first reported a subtrochanteric AFF in a postmenopausal woman with a 13-year history of continuous alendronate therapy that was treated with TPTD, vitamin D, and calcium. However, thigh pain, presented as an initial symptom, intensified, and the magnetic resonance imaging (MRI) showed the appearance of fractures worsening. TPTD treatment commenced, and 6 months later, a repeat MRI showed decreased edema at the fracture sites with faint cortical bridging. Six other reports had similar findings. In all, 7 case reports with 8 patients and 11 AFFs (7 incomplete, 4 complete) were described. All the patients were elderly Asian or Caucasian females (age ranging from 63 to 84 years). TPTD treatment started at 1 to 11 months after detection of AFF. TPTD treatment started at 1 to 11 months after detection of AFF. TPTD was administered for 3 to 24 months. Daily subcutaneous (SC) injection was the most common form of treatment (4 patients) although weekly injection was used in 1 patient. Combination treatments included intramedullary nailing in acute fracture setting or as a prophylaxis (for 4 AFF cases), prophylactic locking plate (for 2 AFFs), vitamin D and calcium supplementation in 2 patients. Fracture healing was observed after 3 to 21 months after TPTD treatment. One patient was additionally treated with strontium ranelate because of persistent discomfort after 24 months of TPTD treatment. [25-31]

Miyakoshi et al.[32] retrospectively reviewed the medical records of 45 consecutive AFFs in 34 Japanese patients who had received oral BPs (alendronate or risedronate) for osteoporosis before AFF and had been followed for >12 months (range, 12 to 90). Thirty-seven complete or incomplete AFFs (82%) were treated surgically and 8 incomplete AFFs (18%) were treated conservatively. Based on TPTD...
| References                          | No. patients/side | Age/gender | Time until TPTD | Treatment dose | Duration of treatment | Combined treatment | Key result                                                                                                                                                                                                 |
|------------------------------------|-------------------|------------|-----------------|----------------|-----------------------|--------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Case report                         |                   |            |                 |                |                       |                    |                                                                                                                                                                                                          |
| Gomberg et al. (2011) [25]          | 1, both incomplete fracture | 63/F Caucasian | 5 mo            | PTH 1-34 20 μg SC daily | 21 mo                |                       | At 6 mo of treatment, decreased marrow edema and cortical bridging. At 12 mo, decreased pain with increase in markers of bone metabolism. At 16 mo, resolution of pain. At 21 mo, complete healing on magnetic resonance imaging. |
| Carvalho et al. (2011) [26]         | 1, Rt. complete    | 79/F Caucasian | 10 mo           | PTH 1-34 20 μg SC daily | 3 mo IM nailing      |                       | After 1 mo, closure of the right femur fracture. After 3 mo, s-CTX and serum osteocalcin increased by 22% and 300%, respectively.                                                                           |
| Huang et al. (2012) [28]            | 1, Rt. incomplete fracture | 63/F Asian (Taiwanese) | 1 mo            | PTH 1-34 20 μg SC daily | 9 mo, followed by raloxifene Vitamin D, calcium |                       | After 2 mo, the pain during walking diminished. After 3 mo, fracture line less visible. After 9 mo, thigh tenderness disappeared. After 10 mo, walk without support 1 yr after. After 12 mo, return to previous work. After 15 mo, fracture line completely invisible. |
| Lampropoulou-Adamidou et al. (2013) [29] | 1, Lt. complete fracture, Rt. incomplete fracture | 84/F Caucasian | 11 mo           | Unspecified | 24 mo, followed by strontium ranelate IM nailing on Lt. |                       | After 4 mo, complete healing of left femur fracture. After 11 mo, pain on right femoral diaphysis with subtle localized periosteal reaction. After 1 yr of strontium ranelate treatment, complete resolution of symptoms. |
| Tarazona-Santabalbina and Aguilella-Fernández (2013) [30] | 1, Lt. complete | 73/F Caucasian | 2 mo            | PTH 1-34 20 μg SC daily | 12 mo Vitamin D, calcium |                       | Pain settled down to complete absence. The lytic image reduced in size more slowly, but reached full repair                                                                                                 |
| Fukuda et al. (2014) [27]           | 1, Rt. complete, Lt. incomplete fracture | 74/F Asian (Japanese) | 5 mo            | TPTD 56.5 μg wkly | 8 mo IM nailing both femur, low-intensity pulsed ultrasonography |                       | Union of bilateral femurs achieved after 3 mo of a once-weekly administration of TPTD.                                                                                                                        |
| Tsuchie et al. (2015) [31]          | 2, Rt. incomplete, both incomplete fracture | 78/F, 77/F Asian (Japanese) | 1 mo, 6 mo | Not specified | 1 yr, followed by eldecalcitol Locking plate |                       | Fracture line disappeared 3 and 6 mo after surgery. No thigh pain of the after 2 and 3 yr postoperatively.                                                                                           |
| Case series                         | 34                | (44 fractures) Asian |                | PTH 1-34 20 μg daily in 17 |                       | IM nail 13 Locking plate 3 in TPTD group IM nail 20 Locking plate 1 in non-TPTD group | [44 femora: TPTD group 21 (11 complete, 10 incomplete), non-TPTD group 24 (21 complete, 3 incomplete)] In AFF's treated surgically, mean time to fracture healing significantly better in the TPTD group. Frequency of delayed healing or non-union significantly lower in the TPTD group. | (Continued to the next page)
use after fracture, these AFFs were divided into non-TPTD (n = 24) and TPTD (n = 21) treated groups. In subanalyses for all AFFs treated surgically, mean time to fracture healing was significantly better in the TPTD group (5.4 ± 1.5 months) than in the non-TPTD group (8.6 ± 4.7 months, P = 0.012), and the frequency of delayed healing or non-union was significantly lower in the TPTD group than in the non-TPTD group (P = 0.014). On the other hand, subanalyses for incomplete AFFs treated conservatively showed no significant differences between the groups.

| References | No. patients/side | Age/gender | Time until TPTD | Treatment dose | Duration of treatment | Key result |
|------------|-------------------|------------|-----------------|-----------------|----------------------|------------|
| Saleh et al. (2012) [33] | 10 All incomplete (4 bilateral) | 67 ± 10 (range, 50 to 85) | Not specified | IM nail in 7 fractures | Administration of TPTD was associated with an increase in all 3 dynamic histomorphometric parameters (bone formation, mineralizing surface, and mineral apposition). |
| Miller and McCarthy (2015) [34] | 15 | 67 (range, 57 to 87) | 6 wk-7 mo | PTH 1-34 20 μg daily in 17 | Ten fractures without a radiolucent fracture line treated successfully with partial weight-bearing restrictions and TPTD. Of 9 fractures with radiolucent fracture line, 2 were treated successfully with TPTD. Six underwent surgical prophylaxis after 3 mo of TPTD. One underwent surgical prophylaxis without it. |
| Chiang et al. (2013) [23] | 14, 6 complete, 8 incomplete | 76 ± 19, 13/F, 1/M | 20 μg SC daily | 6 mo | In 5 treated patients, 2 to 3 fold increase in bone remodeling markers and fracture healing in treated patients. Two patients became pain-free and the remaining 3 patients had improvement in pain. |

**Table 1.** Continued

TPTD, teriparatide; PTH, parathyroid hormone; SC, subcutaneous; Rt., right; Lt., left; IM, intramedullary; CTX, C-terminal telopeptides of type I collagen; mo, month; yr, year; wk, week; wkly, weekly; F, female; M, male; AFF, atypical femoral fracture.
ing. Of 9 patients managed conservatively or surgically without TPTD, 6 had poor fracture healing with ongoing pain, one sustained a contralateral atypical fracture, and one had fracture union after 1 year (Table 1).[23]

**DISCUSSION**

AFF wreaks catastrophe on osteoporotic patients who have had prolonged BP treatment. The over-mineralized fracture ends in these patients pose challenge to fracture union because of delayed bone remodeling. TPTD, which is a bone-forming agent used for nonunion, has become a key therapeutic agent for promoting bone union in AFF. A recent paper elucidates a working mechanism that can possible explain the action of TPTD in AFFs. Murphy et al. [35] employed a mixture of traceable BPs, namely fluorescent-labeled pamidronate and radiolabeled zoledronic acid in an animal study. The traceable BPs were dosed in Wistar rats for models of normal growth and closed fracture repair. Consistent with increased BP remobilization with PTH (1-34) (a form of TPTD), there was a significant decrease in fluorescence in both the long bones and in the fracture callus in PTH-treated animals compared with controls rats. Increased intracellular BP was noted in resorbing osteoclasts, confirming that, in principle, PTH (1-34) increased bone turnover as well as BP turnover.

Although there is a lack of level 1 study on the evidence of TPTD in promoting bone union in AFFs, our systemic review of the literature reveals that TPTD works positively for AFFs. Case reports continue to provide anecdotal evidence of the efficacy of PTH for fracture healing but are of limited utility in guiding clinical decision making. The prospective study of Chiang et al.[23] shows the advantages of using TPTD in AFFs that have inherent difficulty in achieving fracture healing. The results from the study of Miyakoshi et al. [32] show that TPTD administration shortens the union time and reduces the incidence of delayed or nonunion in surgically treated AFFs while it is undetermined in non-surgical cases. The series of Saleh et al.[33] shows that incomplete fractures without radiolucent lines are responsive to TPTD alone whereas those with radiolucent line needed intramedullary nailing. These results imply that TPTD works best when the fracture site is stable, either inherently or with surgical fixation.

Addressing fracture healing in AFFs is very important, and it can be achieved using anabolic therapy. We have conducted a systematic review of the literature on the use of TPTD to enhance fracture healing. Our study has the limitation of depending on the quality of the studies it analyzed. Due to limited available evidence on TPTD use for AFFs, we relied considerably on case reports and retrospective case series in this study. While anecdotal evidence of beneficial effects of TPTD on fracture healing offer limited guidance for clinical decision making, a better understanding of the role of TPTD in fracture healing may be elucidated with future prospective trials. As such, TPTD is currently a viable treatment option to enhance fracture healing in AFFs.

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