Bisphosphonate-Related Osteonecrosis of the Jaw Bones in a Patient with Ankylosing Spondylitis Receiving Anti-Tumor Necrosis Factor Treatment

Banu İŞLETEN, Simin HEPGÜLER, Selen BAYRAKTAROĞLU, Gökhan KESER

1Department of Physical Medicine and Rehabilitation, Medical Faculty of Ege University, İzmir, Turkey; 2Department of Radiology, Medical Faculty of Ege University, İzmir, Turkey; 3Department of Internal Diseases, Medical Faculty of Ege University, İzmir, Turkey

Bisphosphonate-related osteonecrosis of the jaw bones is a rare, but well-recognized pathology, occurring mainly in patients receiving parenteral and high doses of bisphosphonates for the treatment of skeletal metastasis and/or hypercalcemia associated with cancer. However, to a lesser extent, this complication may also occur in patients receiving oral bisphosphonates for the treatment of osteoporosis. In this article, we present a 58-year-old female patient with ankylosing spondylitis (AS) who developed mandible osteonecrosis following long-term oral alendronate treatment for osteoporosis. Dental tooth extraction possibly triggered the occurrence of mandible osteonecrosis in this patient. This patient is notable for receiving concomitant anti-tumor necrosis factor (anti-TNF) treatment. To our knowledge, occurrence of bisphosphonate-related jaw osteonecrosis in a patient with AS receiving concomitant anti-TNF treatment has not been reported previously in the literature.

Key words: Ankylosing spondylitis; bisphosphonate; jaw osteonecrosis.

Bisphosphonates, the potent inhibitors of osteoclast-mediated bone resorption, are mainly used in the treatment of metabolic and oncological diseases involving the skeleton, including osteoporosis, Paget’s disease, and metastatic bone lesions.[1] They have a high affinity for hydroxyapatite bone mineral surfaces, especially in regions with high bone-remodeling. A subgroup of these agents known as aminobisphosphonates, which includes alendronate, also inhibits the osteoclastic enzyme known as farnesyl pyrophosphate (FPP) synthase, thereby maximizing their antiresorptive potential.[2]
Oral bisphosphonates used for the treatment of osteoporosis are generally well tolerated with some predictable side effects, such as gastrointestinal intolerance. However, in recent years, bisphosphonate-related osteonecrosis of the jaw (ONJ) has also appeared in the literature as a rare but serious side effect of bisphosphonate therapy, although a direct causative relationship has not been demonstrated.

Ruggiero et al. [3] reported this complication first in 2003. Osteonecrosis of the jaw is currently defined as an area of exposed bone in the maxillofacial region that has not healed within eight weeks after identification by a healthcare provider in a patient who is receiving or has been exposed to a bisphosphonate and has not had radiation therapy to the craniofacial region. [4] Most of the reported cases are cancer patients receiving parenteral high doses of bisphosphonates, with osteoporosis patients receiving oral bisphosphonates being the next highest group.

Hereby, we present a female with ankylosing spondylitis (AS) who developed mandible osteonecrosis after long-term oral alendronate treatment for osteoporosis. Dental tooth extraction preceded the occurrence of mandible osteonecrosis in this patient, who is also notable for receiving concomitant anti-tumor necrosis factor (anti-TNF) treatment for three years.

**CASE REPORT**

In September 2008, a 58-year-old female diagnosed with AS in 1973 was referred to Ege University Hospital, Department of Physical Medicine and Rehabilitation complaining of pain and discomfort in the right mandible. These symptoms had started in May 2008 following a tooth extraction. The extraction socket did not heal, even after eight weeks, despite antibiotic therapy. Besides the pain and discomfort, she experienced the presence of some particles in the tooth extraction socket of her right mandible which were probably autogenous bone. A panoramic radiograph was taken 10 weeks later in July 2008 which revealed irregularity and diffuse sclerotic bone changes in the trabecular bone structure of the right mandible as well as a bony defect at the level of the extraction socket (Figure 1). A sample obtained from the exposed area was sent for culture to the microbiology laboratory due to the suspicion of dental infection/abscess. Oral flora microorganisms grew in the culture.

Her past medical history related with AS treatment documented that she had received primarily indomethacin between 1973 and 1996. In addition, during her peripheral arthritis attacks, she had intermittently received prednisolone starting with 20 mg/day with doses tapering and completely stopping after a few months. In 1988, she had surgery for a total hip replacement.

In 1996, methotrexate (MTX) and sulphasalazine (SSZ) were commenced. However, MTX was discontinued due to hepatic side effects. Because of the clinical disease progression and activity, anti-TNF treatment (etanercept 25 mg twice weekly) was started in April 2004. A tuberculin skin test was negative, so latent tuberculosis (TB) prophylaxis was not given. She benefited from etanercept and remained on this treatment for more than three years until September 2008.

She received bisphosphonate treatment continuously for 10 years between 1998 and 2008 because of osteoporosis. Oral alendronate had been used as 10 mg/day in the first five years, and when once-weekly doses became available, she received 70 mg/week for the remaining five years. Although past bone densitometry reports were not available, current t scores were -3.2 in the lumbar region and -1.5 in the femoral regions.

A recent physical examination revealed that besides the mandible problems, the patient’s blood pressure, pulse, and body temperature along with her cardiac, respiratory, and abdominal systems were within normal limits. In a locomotor system examination, cervical and lumbar ranges of motions were quite restricted. Tragus-wall distance was 18 cm, lombar flexion 0 cm, cervical rotation 0 cm, lumber lateral flexion 5 cm, and intermalleolar distance 20 cm. The Bath Ankylosing Spondylitis Disease Index (BASDI) and Bath Ankylosing Spondylitis Functional Activity Index (BASFI) scores were 4.74 and 9.21, respectively.

![Figure 1. Panoramic radiograph of the patient showing diffuse sclerotic bone changes posterior to the right mandible around the first premolar teeth (short white arrows). There is a bony defect at the level of the extraction socket (long white arrow).](image-url)
Besides a mild elevation in acute phase reactants, routine biochemistry and full blood count examinations were within normal limits.

Since she had had a history of etanercept treatment for three years, an autoantibody profile was also ordered. She was found to have positive speckled antinuclear antibody (ANA) with a 1/40 titer; other autoantibodies including antiphospholipid (aPL) antibodies were negative. The autoantibody profile was not available before the etanercept treatment began.

Together with the past history of long-term alendronate treatment and occurrence after tooth extraction, relevant clinical symptoms and conventional radiographic findings led to the suspicion of ONJ in this patient. We immediately ordered dental computed tomography (CT) which revealed sclerosis and osteolytic changes in the right mandible. This confirmed the diagnosis of osteonecrosis (Figures 2, 3). The patient also consulted with dental surgeons, and since there was no pathologic fracture, extra-oral fistulae, or osteolysis extending to the inferior border of the mandible, the disease was accepted to be in the second stage. Surgery was not recommended, and conservative management was started. The patient was advised to discontinue the alendronate treatment, and the etanercept treatment was also stopped. A good oral hygiene regimen using antibacterial tooth rinse was recommended. Her complaints and symptoms alleviated gradually. A control panoramic radiography taken after one year of bisphosphonate withdrawal displayed healing of the exposed bone (Figure 4).

Currently, she is receiving only indomethacin supp 100 mg/day for AS along with 600 mg/day calcium and 400 IU/day vitamin D for osteoporosis.

**DISCUSSION**

The present case report describes the occurrence of ONJ in an elderly woman with AS who received long-term oral alendronate for osteoporosis treatment. This patient had had a history of concomitant etanercept treatment for more than three years. Unlike high doses of bisphosphonates given by the intravenous route, as in the case of cancer patients, the prevalence of ONJ due to oral bisphosphonates for osteoporosis treatment is very low and reported to be 0.00038%. To our knowledge, the occurrence of this very rare complication in a patient with AS receiving concomitant etanercept treatment has not been previously reported in the literature.
In the present case, the diagnosis of ONJ was made based on the presence of relevant clinical and radiological findings as well as the past history of oral alendronate treatment for 10 years and occurrence after tooth extraction. These features are consistent with the data in the literature documenting that being an older female and having long term bisphosphonate treatment (generally longer than 3 years) along with previous invasive dental treatment were the most common characteristics of the previously reported cases who developed ONJ during osteoporosis treatment.[6,7] By collaborating with dental surgeons, we made a detailed differential diagnosis in this patient and excluded other possibilities, such as periodontitis, osteomyelitis, periapical disease, gingivitis, tumors or metastasis, and temporomandibular joint disease.

Although the exact mechanism of ONJ remains unknown, suppression of bone remodeling and angiogenesis as well as toxic effects due to oral epithelium seem to be responsible.[8,9] Bisphosphonates substantially reduce bone turnover, impairing the repair of microdamage. They also cause increased bone mineralization which increases the bone brittleness. The microdamage and microfractures that occur either physiologically as the result of the constant stress of masticatory forces or pathologically as the result of local infections and dental extractions cannot be repaired. The impaired bone architecture and quality increases bone fragility, finally leading to the development of ONJ.[8,9] Infections generally accompany ONJ, which not only increases the tendency for ONJ development, but also exacerbates the clinical symptoms. Likewise, it is well known that ONJ may be asymptomatic in early stages. These symptoms mostly occur due to secondary infections,[9,10] as was observed in the present case. Osteonecrosis of the jaw preferentially involves the bones that have a high intracortical remodeling rate, which explains why it occurs mostly in the mandible (70%) and less frequently in the maxilla (30%).[9,10]

The development of ONJ has not been known to be directly caused by the presence of AS nor by treatment with etanercept. Dental tooth extraction seemed to be the main triggering factor for ONJ in our patient. Comorbid diseases or other concomitant medications should also be considered as potential precipitating factors; however, our case did not have any comorbid diseases. With respect to concomitant medications, corticosteroids and immunosuppressive agents are reported to contribute to osteonecrosis development.[11] Therefore, the past history of intermittent corticosteroid use in our patient should also be kept in mind. Besides, thrombogenic aPL antibodies may also precipitate osteonecrosis;[12] however, our patient was negative for these antibodies.

To our knowledge, ONJ should not be included as one of the complications of anti-TNF agents. In our case, etanercept treatment cannot be hold responsible from the occurrence of the ONJ complication. If anti-TNF agents really facilitated ONJ development, this effect could not be explained solely by causing a tendency toward infection. Anti-TNF agents are well known to cause a modest increase in bone formation and suppression in bone resorption.[13] Consequently, this resorption may contribute to ONJ development. Evidence supporting this assumption comes from the fact that anti-RANKL (receptor activator of nuclear factor-kB ligand) antibody (denosumab) treatment for osteoporosis may also induce osteonecrosis development, which is similar to that of bisphosphonates.[14] This may imply that any therapeutic agent which suppresses bone resorption, including etanercept and denosumab, may facilitate osteonecrosis development.

The management of ONJ is initially conservative, including ending the bisphosphonate treatment, eliminating pain, controlling the infection, and preventing the progression of osteonecrosis.[8,10] A good oral hygiene regimen with oral antimicrobial rinses is necessary. Identifying the stage of ONJ is also important for determining proper management. Osteonecrosis of the jaw is classified as having stages from 0 to 3 according to clinical signs and symptoms.[10] No imaging criteria are used for this classification. Delaying surgical interventions until stage 3 is recommended. If surgery becomes necessary, the necrotic area should be extracted with minimal trauma to surrounding healthy tissues.[10] Our patient was accepted as stage 2 and a conservative approach proved to be sufficient with no surgical treatment needed.

Although the risk of ONJ development is low during osteoporosis treatment, preventive measures should always be considered. Before starting the treatment, a dentist should be consulted and, if needed, should initiate it. It is also recommended that the bisphosphonate treatment be stopped temporarily before a dental surgery. It should only be restarted after complete healing of the bone, especially for patients who received bisphosphonate for more than three years.[8]

In conclusion, ONJ development during oral bisphosphonate treatment in osteoporosis patients is
a very rare, but well-recognized complication. To our knowledge, occurrence of this very rare complication in an osteoporotic patient with AS has not been reported in literature. Although, dental tooth extraction along with long term alendronate treatment seem to be the main causative factors for this patient, the possibility of even a minor contribution of etanercept by means of facilitating oral infection and suppressing bone resorption is speculative. Osteoporosis may commonly accompany AS, and bisphosphonate treatment has a potential to be used in AS, both for disease control and OP treatment. As the use of bisphosphonates in AS patients increases, the incidence of ONJ may also be expected to rise in the future.

Declaration of conflicting interests
The authors declared no conflicts of interest with respect to the authorship and/or publication of this article.

Funding
The authors received no financial support for the research and/or authorship of this article.

REFERENCES