Osteonecrosis of the Jaw (ONJ) mainly associated with metastatic breast cancer and multiple myeloma taking the bisphosphonates [2]. Initial reports showed that ONJ was less commonly ONJ appears to occur spontaneously in patients diagnosed after invasive dental procedures such as tooth extraction. Of reported cases of bisphosphonates associated with ONJ have been drugs, corticosteroids, and then also bisphosphonates. The majority (ONJ) associated with bisphosphonate therapy in cancer patients. These density will increase even though the bone volume does not. Bisphosphonates may also contribute to an inhibition of bone resorption and an increase in bone mass. Although the bisphosphonate drugs block bone resorption, bone formation continues for about six to twelve months, after which formation then stops. Thus the newly-mineralized bone is more densely packed so that individual bone density will increase even though the bone volume does not.

Wang et al. [1] reported the first case of osteonecrosis of the jaw (ONJ) associated with bisphosphonate therapy in cancer patients. These cancer patients were undergoing many treatments with chemotherapy drugs, corticosteroids, and then also bisphosphonates. The majority of reported cases of bisphosphonates associated with ONJ have been diagnosed after invasive dental procedures such as tooth extraction. Less commonly ONJ appears to occur spontaneously in patients taking the bisphosphonates [2]. Initial reports showed that ONJ was mainly associated with metastatic breast cancer and multiple myeloma [3] and it was clear that patients taking higher cumulative doses of bisphosphonates had greater risk of developing ONJ. ONJ is more prevalent in patients with malignancies than patients with benign bone disorders [4].

Further, Hoff et al. suggested that the fact should not be overlooked that bisphosphonates offer major therapeutic benefits to individuals with metastatic and metabolic bone disease [4]. Bisphosphonate therapy should remain an important medical treatment yet there are a number of implications for dentistry. Dental therapy for the increasing numbers of aging women being prescribed bisphosphonates in treatment for post-menopausal osteoporosis should focus on prevention of ONJ by management of dental disease.

Osteoporosis is a common bone-resorptive, host-dependent, multifactorial and systemic skeletal disease generally affecting older females, marked by reduced bone strength, and decreased bone mineral density and microarchitectural deterioration of bone tissue, resulting in increased bone fragility and risk of bone fracture [5]. Dental implants meanwhile are increasingly sought out by the same aging population. It is clinically necessary to understand the effect on dental implant success of skeletal low bone mineral density and treatment for osteoporosis with oral bisphosphonates. Most investigations conclude that no compelling theoretical or practical basis exists to expect osteoporosis to be a risk factor for osseointegrated dental implants [6-9].

It is estimated that more than five million people are annually treated with oral bisphosphonates for osteoporosis [10]. Patients taking the oral bisphosphonates have a lower incidence of ONJ [11]. In at-risk populations alendronate and risedronate (Actonel®, Warner Chilcott Laboratories) are potent second-generation oral bisphosphonates which can reduce the risk of hip and vertebral fracture by as much as fifty percent [12].

In our ongoing research we hypothesized that once-yearly IV bisphosphonate zoledronic acid (Zometa® or Reclast® by Novartis) to prevent osteoporosis in elderly nursing home patients may cause ONJ. So far 252 patients have been screened and followed one year after conclusion of IV bisphosphonate treatment. No osteonecrosis of the jaw has been observed in any case.

Collaborative ongoing research by this author investigating dental implants placed in post-menopausal women has shown that women have the same dental implant survival potential regardless of bisphosphonate use. Follow-up for two years on twenty patients has shown no failing dental implants or ONJ (unpublished data, this author). Likewise oral bisphosphonates did not have the reverse effect on dental implants and we did not observe any osteonecrosis of the jaw. These conclusions, while promising for the clinician placing dental implants, demand further investigations with significantly larger populations over longer periods of bisphosphonate treatment exposure.

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