Bisphosphonates for osteoarthritis prevention: “Holy Grail” or not?

Kenneth G Saag

The search for effective disease modifying osteoarthritis (OA) agents has been a long and winding road with much unrealised promise. While a debate persists about whether OA is predominately a disease of bone, cartilage, or both, there has been interest in the possible OA disease modifying effects of bone antiresorptive agents including oestrogens, selective oestrogen receptor modulators, calcitonin and bisphosphonates. Bisphosphonates, the most commonly used therapeutic agents for osteoporosis prevention and treatment, hold OA treatment appeal based on their known pharmacology of altering bone remodelling through a direct inhibitory effect on the osteoclast. Such effects could retard subchondral bone remodelling, believed important by some in osteophyte formation and subchondral sclerosis. Also of potential benefit in OA, bisphosphonates have effects beyond the osteoclast and may be slightly immunomodulating via inhibition of pro-inflammatory cytokines. In vitro, etidronate binds to human cartilage and can modestly inhibit matrix metalloproteinases. Neridronate stimulated osteoblasts to produce osteocalcin in an OA model, providing yet another potential pathway for bisphosphonate activity in OA. Animal data supports a possible reduction in osteophyte progression and a suppression in subchondral bone resorption with bisphosphonates. Rabbits treated with zoledronic acid and rats receiving alendronate experienced chondroprotection. Bisphosphonates may also stimulate collagen and proteoglycan synthesis.

Carbone and colleagues examined women in the Health, Aging, and Body Composition Study, observing a reduction in magnetic resonance imagery (MRI)-associated marrow oedema in subchondral bone but no reduction in knee OA radiographic progression among those receiving alendronate. Cohort studies of this type are limited potentially by systematic differences in OA predilection among those who do and do not receive bisphosphonates. Randomised controlled trials of risedronate initially suggested a bisphosphonate OA benefit in terms of pain, function, radiographic joint space changes and retained vertical trabecular structure. Spector and colleagues found a reduction in the Western Ontario and MacMaster Universities (WOMAC) OA index, along with a non-significant trend towards reduced joint space narrowing in patients receiving three times the usual osteoporosis dose of risedronate (n = 284 patients). However, a subsequent much larger randomised controlled trial (n = 2483 patients) did not demonstrate a clear OA advantage with respect to radiographic joint space narrowing or WOMAC subscales; however, it was observed that the biochemical measure of cross-linking telopeptide of type II collagen was reduced compared to placebo. A subset analysis of the study population who were “rapid joint space losers” and who were randomised to risedronate experienced more retained vertebral trabecular structure (particularly at a 50 mg/week risedronate dose) than those receiving placebo. Some have speculated that this larger clinical trial was underpowered based on a low rate of OA progression in the placebo group. Further, the timing of potential benefit of antiresorptives in OA may be critical with the greatest benefit seen very early in the disease course, although longer-term follow-up for late effects is also missing in many clinical trials. Lastly, it should be noted that independent of their effects on spinal fractures, several bisphosphonates have reduced low back pain. This finding raises questions about whether this effect alternatively could be mediated by a reduction in spinal OA progression.

In this issue of Annals of the Rheumatic Diseases, Neogi and colleagues (see page 1427) report on aycler conducted secondary analysis of 200 patients randomly selected from the large and well known Fracture Intervention Trial (FIT). When radiographs were carefully read against a standard atlas they found a small reduction in spinal osteophyte progression in those patients randomised to receive alendronate compared to placebo. While the spinal radiographic changes were significant, they were subtle and the authors acknowledged that this level of radiographic difference may be of minimal clinical relevance. Similar benefits were seen in terms of disc space narrowing in the lumbar spine, although the significance of this finding did not persist when the thoracic spine was included in the analysis. Although this secondary analysis of FIT was very carefully conducted, there was only modest correlation between the radiograph readers, further questioning the clinical relevance of the findings. Overall, the radiographic reading methods used should have resulted in a non-differential outcome misclassification between those receiving and not receiving alendronate. This study was limited by the fact that the results may not generalise to those without osteoporosis. Since OA occurs preferentially among those without osteoporosis, most studies of OA have been conducted in these patients.

Does this paper support a stronger role for bone in the pathogenesis of osteoarthritis or for bisphosphonates as agents that may retard OA progression? Are the effects of bisphosphonates differential and, if so, could the alendronate used in this study be more efficacious than the very equivocal (likely negative) evidence supporting risedronate as an OA therapy? These and other important clinical questions cannot be fully answered by this paper or by the existing literature, but the results from this important study by Neogi and colleagues provide clear further rationale for additional large-scale comparative clinical trials of bisphosphonates and other bone antiresorptive agents in OA. Since many osteoporosis studies obtain serial spinal radiographs, it would be prudent that this information be analysed as a possible osteoarthritis endpoint. Even if it can be much more consistently demonstrated that all or certain bisphosphonates reduce OA progression in the spine, it is not at all clear that this will correlate with benefits at the knee or hip; the key anatomic areas that lead to the most OA morbidity. In addition to the need for these further studies evaluating bisphosphonate efficacy OA, a recent spate of mostly uncontrolled observations suggest possible new bisphosphonate safety issues.
One such potential safety signal of exposure to bisphosphonates with long skeletal retention is over-suppression of bone turnover leading to a proposed increased rate of atypical insufficiency fractures. So far, the evidence justifying this and other newer safety signals such as atrial fibrillation is rather limited, but additional pharmacoepidemiological studies on these topics will allow for eventual better determination of a bisphosphonate benefit–risk ratio. Pending these now more urgently needed further studies on OA efficacy and longer-term safety with bisphosphonate, when an osteoporosis therapy is otherwise needed, a potential beneficial effect on OA might provide slight added value in the decision to prescribe a bisphosphonate.

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