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To the Editor:

As a rheumatologist who has seen and managed bisphosphonate related osteonecrosis of the jaws (Fung E, Medwatch case reports; Harper and Fung3), I wish to comment and raise questions on the recent editorial by Kahn2 and article by Etmiman, et al3.

Osteonecrosis of jaws (ONJ) might be old, dating back a century in the form of "phossy jaw"; but bisphosphonate related ONJ is rather new, first reported in 20033. By the current, mostly accepted, definition, it is certainly not equal to osteoradionecrosis and we should not confuse the 2 conditions.

The incidence of bisphosphonate related ONJ in oral therapy was estimated as 0.7 or < 1 per 100,000 patient-years5; it is not the < 1 per 100,000 patients that has been so frequently misquoted2. The importance of getting it straight is, as rightfully pointed out by Dr. Kahn2, that the duration of therapy might be a very important factor, particularly in view of the very long bone half-life of amino-bisphosphonates6.

In the editorial2, the work by Mavrokokki, et al from Australia in 20077 is not mentioned. I remind readers that that work cannot be taken lightly, in view of the country-wide design of the survey, which was supported by the national healthcare service and also by the centralized electronic records for dental care used in South Australia. The results for bisphosphonate related ONJ, as published, for osteoporosis treated with oral bisphosphonate were as high as 1/2260 cases without and 1/296 cases with dental extraction. While these findings seemingly are at odds with the North American experience5 (except the recently reported prevalence of bisphosphonate related ONJ of 1/1424 cases from Northern California8), the incidence for malignancy treated with bisphosphonate of 1.15% without and 9.1% with dental extraction is in agreement with data reported in North America9,10.

The article by Etmiman, et al3 shows a 3-fold increase of aseptic osteonecrosis with bisphosphonate, but the site of the aseptic osteonecrosis cannot be verified. In fact it might be at sites other than the jaw, since there was no International Classification of Disease code for it prior to November 2007, and Etmiman, et al felt these findings might be corticosteroid related in view of the imbalance of the percentages of corticosteroid use. However, within their Methods section, Etmiman, et al stated that potential confounders including oral corticosteroids were adjusted for by conditional logistic regression. Did Dr. Etmiman actually adjust or not adjust for the corticosteroids in this study? This might fall short compared to the Australian report, in which cases of ONJ were confirmed, although not adjudicated, by experienced clinicians.

Finally, the only bisphosphonate clinical study that has included ONJ in the protocol is the zoledronic acid Horizon trial11. If indeed the incidence is as high as the Australian survey indicates, we could have seen some signals of it; on the other hand, if it is anywhere close to the finding of < 1 per 100,000 patient-years, then, as Etmiman and colleagues commented, none of the bisphosphonate trials to date has the statistical power to reveal this one way or the other.

Bisphosphonate related ONJ does not discredit bisphosphonates as an effective osteoporosis therapy, but our patients deserve more than having their treating physicians look the other way from this new development, or entity. Adding to the laborious Australian survey7 and the thoughtful American Society of Oral and Maxillofacial Surgeons 2007 position statement12, we should welcome the results of the Canadian survey3 as well as the authors’ position in this important issue.

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