Uncoupling of disease activity and structural damage. Does it matter clinically?

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Rheumatoid arthritis (RA) is an autoimmune disease characterised by synovial inflammation that can lead to joint damage through bone and cartilage destruction, loss of function and decreased quality of life. Fortunately, over the past few years, we have witnessed major advances in the management of RA with new agents and treatment strategies that have improved outcomes and made remission become a realistic goal. True remission has been defined as a state where there is no evidence of disease activity with complete resolution of signs and symptoms as well as arrest of joint damage and disability progression. Hence, inhibition of radiographic progression has become a major therapeutic goal given the potential for prevention of destruction, loss of function and decreased quality of life.

Clinical remission can be achieved with disease-modifying antirheumatic drugs (DMARD) alone or may need a combination of DMARDs and biologic agents. Radiographic remission is more complex as we have learned from the probability plots that the mean/median change of radiographic progression is accounted for by a minority of patients who progress radiographically. Indeed, regardless of the therapy used, approximately 50% of patients will not show radiographic progression over 2–3 years. This percentage increases up to 75–85% of patients receiving combination therapy, and those who do progress, progress less than patients on methotrexate alone.3

It has long been thought that prevention of radiographic progression was achieved through the eradication of synovial inflammation, and that antitumour necrosis factor (TNF) agents were more effective at reducing structural progression through a better control of inflammation. However, recent posthoc analyses have shown a disconnect between clinical and radiographic outcomes where inhibition of radiographic progression was achieved even in patients with significant residual disease activity treated with TNF inhibitors suggesting a separate direct role for TNF on bone damage.4–5 Interestingly, Aletaha and colleagues have demonstrated a similar dissociation between clinical and structural outcomes in patients treated with rituximab suggesting that the mechanism explaining this dissociation is not unique to anti-TNF drugs.6

The Anti-TNF-α Trial in Rheumatoid Arthritis with Concomitant Therapy (ATTRACT) was the first study to suggest the notion of disconnect, or uncoupling, between damage and inflammation when they showed a reduction in total sharp score in RA patients treated with a combination of infliximab and methotrexate who showed no improvement in tender and swollen joint counts, health assessment questionnaire (HAQ) score, patient global score and C-reactive protein (CRP) compared with patients on methotrexate and placebo.5 Similar findings have been shown in early RA and with other anti-TNF agents.5–7 This suggests that there may be a requirement for a higher TNF level threshold for induction of joint damage than for signs and symptoms, or that the biological processes leading to symptoms are not the same and less driven by TNF than those causing radiographic progression. TNF is thought to cause erosion mainly by its ability to promote the synthesis of receptor activator of nuclear factor kappa-B ligand (RANKL) which, in turn, activates the maturation of osteoclasts that leads to bone erosions. It has indeed been shown that denosumab, an anti-RANKL monoclonal antibody, reduces structural damage in RA while having no effect on clinical symptoms.9 More recently, a similar effect was demonstrated with tocilizumab which inhibits IL-6, a cytokine involved in the activation of osteoclasts.10 It is possible that the level of TNF or IL-6 required to induce joint damage is greater than the one needed to promote clinically detectable synovitis.

However, the explanation may be more complex as the dissociation between clinical and radiographic outcomes also seems to occur when targeting upstream mechanisms such as B-cell depletion with rituximab. Using data from the A Study to Evaluate Rituximab in Combination With Methotrexate in Methotrexate-Naïve Patients With Active Rheumatoid Arthritis (IMAGE) trial, Aletaha et al have shown that the combination of rituximab and methotrexate retards joint damage independently of disease activity, while methotrexate retards joint damage only in patients who had a good clinical response.6 An interesting finding that was not looked for in many of the studies using TNF inhibitors is that this effect was true for both the erosion and the joint space narrowing score, and was consistent when they matched patients for disease activity. As rituximab targets B-cells which are at the beginning of the inflammatory cascade leading to the development of RA, it seems harder to explain why it does not produce the same effect on inflammation and bone damage.

In order to explain this dissociation, Watson et al proposed a ‘two-compartment model’ with centres of pathology in both bone and synovium.11 Autoactive B cells escape deletion in the bone marrow and migrate to either the synovium leading to synovitis or to subchondral bone where they will lead to osteitis and, eventually, bone erosion. Indeed, imaging studies have shown that the best predictor of radiographic progression is not MRI synovitis but bone marrow oedema which correlates with osteitis.12 Once initiated, it could be that the biologic processes occurring at each site are different with predominance for B cells in the bone, and possibly more T cells in the synovium explaining the effect of rituximab on damage even in patients without clinical improvement. Supporting this hypothesis, a study looking at the cellular components of bone oedema, known to be an important predictor of bone erosion, found that these were made predominantly of osteoclasts with a trend for more plasma cells and aggregates of B cells.13 On the opposite side, in a small study examining synovial tissue from five RA patients, 75–90% of lymphocytes present were identified as T cells.14 When comparing the cellular composition of bone marrow aggregates with synovial tissue, Jimenez-Boj and colleagues have also demonstrated...
a higher number of B cells in the bone marrow aggregates and similar number of T cells.15

Finally, these novel findings are mainly interesting because of their clinical implications. As recommend by an expert task force, most rheumatologists now treat their patients to a target which should be remission or low disease activity.16 These recommendations, as well others developed by the different national organizations,17 18 should definitely be implemented in all patients, especially those treated with methotrexate (MTX) or the traditional DMARDs, who display more radiographic damage despite good clinical control.19 However, results from this trial, and others with TNF inhibitors, suggest that this target may not have to be the same for patients on biologics than if on methotrexate only. However, we should still aim at suppression of clinical symptoms as they are what affect patients most on a daily basis, and change treatment in patients with suboptimal clinical response despite the potential prevention of bone damage.

However, several issues remain; the majority of the recent clinical trials, including this one, have demonstrated very little radiographic progression.6 20 Should we start using more sensitive techniques to evaluate structural damage, such as ultrasound, computerised tomography or MRI which will not only assess the structural damage but also residual inflammation in the synovium and bone that can lead to cartilage and bone lesions? Should these new imaging modalities be incorporated in the definition of remission? We would then need to define the threshold of imaging activity that will be clinically relevant.

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