Combination Therapy with Methotrexate and Etanercept for Refractory Chronic Recurrent Multifocal Osteomyelitis

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J Rheumatol 2011;38;782-783
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To the Editor:

Chronic recurrent multifocal osteomyelitis (CRMO) is a form of chronic nonbacterial osteitis (CNO) characterized by one or more lytic bone lesions with no identifiable cause. The metaphyses of long bones and the clavicle are most frequently affected, although any bone may be involved. Most cases begin in childhood, and follow an intermittent course. Pain is the most common presenting symptom. Affected individuals are sometimes febrile, and acute-phase reactants may be increased, pointing to the inflammatory expression of this condition. Lesions may resolve at or before the time of puberty, but in some cases skeletal deformity and/or disability may occur.

We describe 2 children with CRMO whose symptoms persisted after trials of nonsteroidal antiinflammatory medication, glucocorticoids, and methotrexate (MTX). Combination therapy with MTX and etanercept resulted in amelioration of pain, accompanied in one patient by radiographic improvement in a lytic bone lesion.

Case 1. A 4-year-old boy developed pain in his right foot and refused to walk for 2 months. Plain radiographs showed normal-appearing vertebrae. A bone scan disclosed increased uptake in the right talus and navicular bone. He received several courses of intravenous and oral antimicrobials, without improvement. Fever was absent, C-reactive protein was normal, and erythrocyte sedimentation rate (ESR) was mildly increased to 23 mm/h. A second bone scan performed 2 months later showed increased radionuclide uptake in the L4 and L5 vertebrae. Focal magnetic resonance imaging (MRI) revealed vertebral abnormalities consistent with CNO including increased signal intensity of the vertebral marrow (Figure 1A). Talar biopsy was nonrevealing, showing bone fragments with myxoid-appearing fibrous stroma. Naproxen 10 mg/kg/day was given, without improvement. Prednisone 2 mg/kg/day was added for 3 weeks, with reduction in pain, but symptoms recurred during taper. During the third month, oral MTX (15 mg/m² once weekly) was given with naproxen. However, he complained of worsening leg and back pain (verbally described as 7 on a scale of 1–10) and was unable to walk. Etanercept 0.8 mg/kg/week was prescribed with MTX and naproxen. One month later, his limp had resolved completely, and he reported no pain. ESR was 16 mm/h. MRI performed after 6 months of treatment showed diminished signal abnormality, indicating improvement in bone inflammation (Figure 1B), and he remained symptom-free. His care was transferred to another medical center.

Case 2. A 12-year-old girl presented with elbow, ankle, hip, and foot pain worsened by activity for 6 months, without fever. Radiographs disclosed numerous lucent bony lesions with sclerotic margins in the metaphyses of the left distal tibia and fibula, and a healing fracture in the distal portion of the right fourth metatarsal. Focal MRI showed lesions with enhanced T2 signal including pelvic, right ankle, and left foot lesions, as well as in the acetabular roof, pubic rami, second sacral segment, distal tibia and fibula, and fourth and fifth metatarsal bones. Direct biopsy was not performed at request of the family. However, the clinical and radiographic findings were compatible with presumed CRMO. A bone marrow aspirate showed normal cytology. Naproxen 10 mg/kg/day, prednisone (2 mg/kg/day for 1 month, followed by slow taper), and oral MTX (10 mg/m² once weekly) were given in succession. However, after 8 months, she complained of right buttock, hip, and left ankle pain (described as 4 on a scale of 1–10) and limp. Etanercept (0.4 mg/kg/dose subcutaneously twice weekly) was added to MTX. Within several weeks, her pain resolved completely, and ESR declined from 18 to 8 mm/h. Pain did not recur during 15 months of treatment. Because the patient felt well, her family declined followup MRI studies.

We describe 2 children with CRMO who continued to experience pain despite antiinflammatory treatment using naproxen, glucocorticoids, and MTX. However, coadministration of MTX and etanercept, a tumor necrosis factor-α (TNF-α) blocker, resulted in progressive and sustained improvement in pain. In one child, clinical improvement was associated with radiographic improvement in a bone lesion. A limitation of our report is that in accord with the parents‘ wishes, posttreatment MRI was not performed until several months after therapy with etanercept was started. Since the natural history of CRMO may include spontaneous remission, we cannot exclude the possibility that MRI changes over this interval may have been incidental rather than due to etanercept treatment.

Up to 25% of cases of CRMO are accompanied by inflammation involving other organs, including the skin or bowel. Histologic examination of bone lesions may reveal neutrophilic inflammation with increased osteoclasts. These observations suggest that CRMO may be an autoimmune inflammatory disorder. Osteolytic lesions resembling those of CRMO are seen in infants genetically deficient in expression of a natural soluble interleukin 1 receptor antagonist (DIRA). However, unlike CRMO, infants affected with DIRA also exhibit other skeletal abnormalities including generalized osteopenia, suggesting the mechanisms underlying bone pathology in these 2 disorders may not be identical. Majeed syndrome, a syndromic form of CRMO associated with dyserythropoietic anemia, is caused by mutations in LPIN2.

There is evidence for a possible role of the proinflammatory cytokine TNF-α in the pathogenesis of CNO. In a large registry, serum concentrations of TNF-α were increased in two-thirds of patients with active disease. Increased TNF-α expression was observed in biopsies obtained from osteolytic lesions of 2 adults with SAPHO (synovitis, acne, pustulosis, hyperostosis, and osteitis) syndrome, a disorder that presents mainly in adulthood and produces bony lesions similar to those of CRMO. TNF-α promotes osteoclast activation in vitro, representing a possible mechanism by which local TNF-α overexpression may contribute to osteolysis.

Etanercept is a recombinant, humanized dimeric protein containing the extracellular domain of the human p75 TNF receptor fused to the Fc region.
of IgGl, which antagonizes soluble TNF-α. Etanercept was given with MTX in our cases, based on favorable experience using combined therapy with these drugs in juvenile idiopathic arthritis. There are several reports of successful use of infliximab, a chimeric anti-TNF-α antibody, in CRMO and SAPHO, one report on the use of etanercept in SAPHO, and one report of use of the anti-TNF-α antibody adalimumab to treat CRMO. To our knowledge, this is the first description of successful use of etanercept with MTX in children with CRMO.

Bisphosphonates have also been used successfully for cases of CRMO that did not respond to initial therapy. We chose etanercept for our patients because of concern regarding the longterm effects of bisphosphonates on the growing skeleton, and reports of osteonecrosis of the jaw associated with their use in adults. Therapy with TNF-α antagonists may be complicated by hypersensitivity reactions or by infection. In addition, recent data have emerged questioning the longterm safety of these agents in children with respect to risk for malignancy. However, the overall risk for cancer in children receiving TNF-α antagonists for rheumatic disease appears to be quite low.

To date, no specific therapy has been universally efficacious for CNO: treatment failures of bisphosphonates have been reported, and incomplete responses to infliximab or waning efficacy over time have been described. Further, since biologically significant differences exist between different TNF-α antagonists, it cannot be assumed that drugs of this class are interchangeable. A multicenter randomized, controlled trial with predetermined outcome measures comparing efficacy of etanercept with MTX, an anti-TNF-α antibody such as infliximab, and a bisphosphate in CRMO/CNO refractory to initial therapy might provide information useful to clinicians caring for children with this condition. At present, we believe that TNF-α antagonists or bisphosphonates are both acceptable treatment alternatives. Whichever class of drug is chosen, we further believe that before treatment is commenced, the known risks and potential benefits should be explained to patients and their families and their consent should be obtained. In cases when treatment with a TNF-α antagonist is contemplated, etanercept with MTX may be preferred to infliximab, since the former regimen can be given at home rather than by intravenous infusion, and immediate adverse reactions are less likely.

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