Approach Sulfasalazine, MTX With Caution in JIA

BY TIMOTHY F. KIRN
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Snowmass, Colo. — Prescribing patterns for sulfasalazine and methotrexate in patients with juvenile idiopathic arthritis need fine tuning, Patience White, M.D., said at a symposium sponsored by the American College of Rheumatology.

The doctor’s concerns: Sulfasalazine needs to be prescribed more cautiously, and methotrexate dosages are often too high.

Although sulfasalazine is commonly prescribed for juvenile idiopathic arthritis (JIA), there are significant risks to weigh against the benefits in this population, Dr. White, chair of pediatric rheumatology at Children’s National Medical Center, Washington, said. Reports have suggested that in late-onset pauciarticular juvenile arthritis, sulfasalazine decreases symptoms by 50% or more. But “I would urge you to use caution here,” Dr. White said. It may not be as effective in other subtypes of juvenile arthritis, and findings from double-blind trials have been mixed on the issue of whether it really is better than placebo.

Most important, 30% of patients with systemic onset JIA can have a febrile, serum-sickness-like reaction while taking sulfasalazine. This is a risk that deserves serious consideration before the drug is started, Dr. White stressed.

Methotrexate is an established second-line agent, yet the dosages used are often too high.

In a recent investigation comparing the safety and efficacy of parenteral methotrexate 15 mg/m² per week with higher dosages for patients with polyarticular-course JIA, researchers found that efficacy plateaued with the lower dose.

All participants had failed to improve while receiving standard dosages of methotrexate (8-12.5 mg/m² per week) (Arthritis Rheum. 2004;50:2191-2201). Dr. White said she considers 10 mg/m² per week the most effective and appropriate dose; when patients do not respond, she tends to try etanercept rather than increase the dose of methotrexate.

Etanercept clearly has a place in treatment, she said. About 74% of polyarticular JIA patients respond to etanercept, and it can reduce the need for prednisone and methotrexate. The biologic has been shown to have no more toxicity in patients as young as 3 years old than it does in adults.

Various combination treatments have been tried in JIA with some success. However, these reports are based on small groups of patients, so combination therapy is not really evidence based, Dr. White said.

She added that sometimes when a JIA patient stops responding to a particular regimen that once was effective, it is not a failure of therapy. It may be that the patient has simply outgrown his or her dosage.

Children can be more susceptible than adults to certain types of toxicity, and these effects can be subtle. NSAIDs, for example, can make some children hyperactive. Naproxen can cause pseudoporphyria.

Because of their effects on growth, systemic corticosteroids should be avoided unless the diagnosis is absolutely certain, she said.

Intraarticular corticosteroid injections, on the other hand, work well in children. Several studies have demonstrated that triamcinolone is the agent of choice, Dr. White added.

WBC Count, Pain Severity Separate Leukemia From JIA

BY TIMOTHY F. KIRN
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Snowmass, Colo. — Elevated WBC count, pain severity are the best indications that a pediatric patient presenting with joint pain has juvenile idiopathic arthritis rather than leukemia, but there are some subtle clinical differences, Patience White, M.D., said at a symposium sponsored by the American College of Rheumatology.

Joint pain in leukemia is usually much more severe than in JIA, and the patient with leukemia may also have bone pain and night pain, said Dr. White, chief of pediatric rheumatology at Children’s National Medical Center, Washington.

In the child with JIA, fever is often high and spiking; fever associated with leukemia tends to be low-grade.

In addition, 90% of patients with systemic JIA will have an associated rash, compared with 10% of leukemia patients.

Still, the two can look clinically similar, and it’s common for patients with leukemia to end up on a potentially harmful trial of corticosteroids. In questionable cases, the white blood cell count is most revealing.

Dr. White noted the case of an 8-year-old boy referred to her for possible diagnosis of systemic onset JIA.

The patient had painful arthritis and a maculopapular rash over the trunk, wrist, and ankle and had been experiencing daily fevers for 4 weeks. He had lost 5% of his body weight.

Testing for infection, including blood and urine cultures and a Lyme disease assay, had been negative.

A course of treatment with an NSAID had been unsuccessful and, most importantly, the white blood cell count at 9,000 cells/µL, was too low to be consistent with a diagnosis of JIA, Dr. White said.

If the patient had JIA, the WBC should have been at least 30,000 cells/µL.

She ordered a bone marrow biopsy rather than initiate a trial of corticosteroid therapy, and the diagnosis of leukemia was confirmed.