Kikuchi-Fujimoto disease (histiocytic necrotizing lymphadenitis): report of a case with other autoimmune manifestations

Tina Mahajan, MD, Richard C. Merriman, MD, and Marvin J. Stone, MD, MACP

Kikuchi-Fujimoto disease (KFD), or histiocytic necrotizing lymphadenitis, is a benign and self-limited disease that mainly affects young women. Patients present with localized lymphadenopathy, fever, and leukopenia in up to half of the cases. KFD can occur in association with systemic lupus erythematosus. We present the case of a patient with KFD and systemic lupus erythematosus, as well as relapsing polychondritis. This patient had persistently low C4 complement levels, so she was evaluated for a genetic defect in complement production and was found to have two “null” C4 alleles. We believe that this may have contributed to the development of her diseases.

A 56-year-old white woman with no significant past medical history was initially seen at Baylor University Medical Center in April 1998. She presented with acute onset of bilateral tender cervical lymphadenopathy with associated malaise, fever to 103°F, and anorexia. She also had severe arthralgias in her hands and knees and had difficulty walking. Through a lymph node biopsy, the diagnosis of histiocytic necrotizing lymphadenitis, or Kikuchi-Fujimoto disease (KFD), was made. She was treated with a course of sulfa drugs for 10 days and started a second course but discontinued it because she broke out in hives. Her symptoms resolved within 2 months.

The patient was well until January 2001, when she began having “tenderness” in the posterior aspect of her hairline and noticed an enlarged right occipital lymph node. She took acetaminophen for the episodes of tenderness, which would come and go and last about 24 hours. By April 2001, she began experiencing fatigue, malaise, and fevers to 104°F and was hospitalized. A lymph node biopsy again showed the classic finding of KFD: necrosis containing abundant karyorrhectic nuclear debris (Figure). The presence of histiocytes was confirmed with the CD68 stain. Bacterial, acid-fast bacillus, and fungal stains and cultures of the specimen showed no growth. She was treated with a prednisone taper, and the findings resolved.

Soon after her hospitalization, the patient was referred to a hematologist/oncologist to determine if she had lymphoma. Bone marrow aspirate and biopsy results were normal. Serum protein electrophoresis and urine immunoelectrophoresis results also were normal. Laboratory work did reveal a positive antinuclear antigen titer (1:160) with a speckled pattern. A diagnosis of systemic lupus erythematosus (SLE) was entertained, and the patient was referred to a rheumatologist.

The rheumatologist confirmed the diagnosis of SLE. The patient had the following clinical findings: arthralgias, linear
ulcers on the gums, alopecia, and normocytic anemia. In addition, she had a positive antinuclear antigen titer and a positive double-stranded DNA test. Her C-reactive protein level was within normal limits, but she had decreased levels of complement. Her C3 level was 50 mg/dL (normal range, 83–177) and her C4 level was <10 mg/dL (normal range, 12–50). C4 allotyping demonstrated two “null” C4 alleles.

The patient was started on hydroxychloroquine 200 mg twice a day. Her arthralgias and mouth ulcers improved. About 9 months after being diagnosed with SLE, however, she began to experience recurrent episodes of redness and pain in her ears and nose. Her rheumatologist diagnosed relapsing polychondritis. The symptoms would last a few days and then resolve on their own without treatment.

As of December 2006, the patient had gone 5 years without recurrence of lymphadenopathy or symptoms of SLE. Her last flare of relapsing polychondritis was 4 years ago.

**DISCUSSION**

KFD, or histiocytic necrotizing lymphadenitis, was originally reported in 1972 in Japan. It has been reported in several countries since then. It occurs most commonly in young women (1) with localized lymphadenopathy, most commonly in the cervical region (2). It is associated with fever and leukopenia in up to 50% of patients (3).

The differential diagnosis of fever and cervical lymphadenopathy is broad and often leads to an extensive workup. Our patient was tested for tuberculosis, Epstein-Barr virus, cytomegalovirus, HIV, toxoplasmosis, and syphilis. In addition, she had a bone marrow examination to check for lymphoma. All of these studies were negative or normal. Lymph node biopsy results did facilitate the diagnosis. The characteristic histology of KFD is single or multiple areas within the lymph node that contain necrosis and histiocytic cellular infiltrate. The capsule of the node may be invaded, and perinodal inflammation is common (4). Cultures and stains for organisms are negative.

KFD is known to occur in conjunction with SLE (Table (5)). Some experts even suggest that KFD is one unusual presentation of SLE. Santana et al did a Medline/LILACS (Latin American and Caribbean Health Sciences) search in 2003 and found 35 reported cases in which KFD and SLE occurred together. In the majority of the cases, SLE was diagnosed either after or at the same time as the KFD (6). In the case of our patient, SLE was diagnosed about the same time as her second episode of KFD. It is interesting to note, however, that during her first episode, she did present with arthralgias. It is not known whether SLE serologies were checked at that time.

No effective treatment has been established for KFD. It is a benign, self-limited disease that resolves in 1 to 4 months. Patients should be monitored, however, since they may subsequently develop SLE or, in unusual circumstances, develop a recurrence of KFD. Recurrences of the latter are uncommon (7).

In cases in which KFD is diagnosed after or at the same time as SLE, corticosteroids are often used for treatment, often along with hydroxychloroquine (8–10). After treatment with prednisone, our patient received hydroxychloroquine once SLE was diagnosed. She did very well. Her symptoms resolved within 1 month of starting the treatment, and she has not relapsed since.

The etiology of KFD is unknown. Certain causative organisms have been proposed. These include Epstein-Barr virus,
were less than those seen in the general population helped to identify or a “flare” of the disease. Knowing that her baseline levels of C4 because many authorities suggest and has done well for the last several years. The presentation of her SLE, and perhaps even to relapsing polyarthritis is also thought to have an autoimmune etiology, and connective tissue, is another problem that our patient developed. Patients most commonly present with unilateral or bilateral ear inflammation with sparing of the noncartilaginous parts of the ear. The next most common presentation is joint involvement, followed by nasal and ocular involvement. Relapsing polychondritis is also thought to have an autoimmune etiology, with autoantibodies attacking the patient’s cartilage. Complement levels may play a role in the constellation of diseases seen in our patient. The complement system is a set of proteins that aids in phagocytosis, chemotaxis, opsonization, and the clearance of immune complexes. There are three different paths in the complement system: the classic pathway, the alternative pathway, and the mannose-binding lectin pathway. Each pathway is activated differently. The classic pathway is activated by binding to immune complexes (19). C4 is an important component of the classic pathway, and so a deficiency in it results in defective immune-complex clearance. Our patient was found to have persistently low C4 levels. Allotyping had demonstrated two “null” alleles at her C4 locus. C4 is involved in the early part of the classical complement pathway. Homozygous complement deficiency, especially of the classical pathway system: the classic pathway, the alternative pathway, and the mannose-binding lectin pathway (18). Each pathway is activated differently. The classic pathway is activated by binding to immune complexes (19). C4 is an important component of the classic pathway, and so a deficiency in it results in defective immune-complex clearance. Our patient was found to have persistently low C4 levels. Allotyping had demonstrated two “null” alleles at her C4 locus. C4 is involved in the early part of the classical complement pathway. Homozygous complement deficiency, especially of the classical pathway, has been strongly associated with the development of some autoimmune disorders, in particular SLE (20). We believe that low C4 levels in our patient perhaps contributed to impaired clearance of immune complexes and may have predisposed her to SLE, KFD (which may have been an unusual presentation of her SLE), and perhaps even to relapsing polychondritis. After her initial treatment with low-dose prednisone and hydroxychloroquine, she gradually became asymptomatic and has done well for the last several years. It was helpful to know that the patient produced less than the expected amount of C4 because many authorities suggest that low C4 levels may suggest either persistent disease activity or a “flare” of the disease. Knowing that her baseline levels were less than those seen in the general population helped to prevent “overtreatment.” Unfortunately, allotyping is no longer performed in any US laboratories.

We feel that this patient’s KFD is probably an unusual manifestation of SLE. In addition, the patient’s atypical course may be related to her inherited chronically low C4 levels. To our knowledge, previous cases of this syndrome were not evaluated for complement deficiency.