Localized Intra-Abdominal Kikuchi-Fujimoto Disease: a Case Report and Review of the Literature

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ABSTRACT
We described a nine-year-old boy with Kikuchi-Fujimoto disease involving localized the intra-abdominal lymph nodes who suffered from persistent fever.

Elevated serum soluble IL2 receptor and urine β2-microglobulin levels in this patient indicated that the clinical findings were due to hypercytokinemia. Abdominal magnetic resonance imaging with gadolinium enhancement revealed multiple enlarged lymph nodes in the periarcic and supramesentric artery regions, which had low signal intensity on short inversion time inversion-recovery imaging. Since it could not be determined if a malignant lymphoma was present from laboratory and imaging studies, we recommended resection of the involved lymph nodes to make a definitive diagnosis and decide whether or not to administer steroids.

Key words ① Kikuchi-Fujimoto disease ② subacute necrotizing lymphadenitis ③ soluble IL2 receptor

Kikuchi-Fujimoto disease (KFD), also known as subacute necrotizing lymphadenitis, is a benign and self-limiting disease with symptoms and signs of fever, eruption, and lymphoadenopathy\(^1,2\). Although KFD was first reported in Japan, it has recently been reported in countries other than ones in East Asia\(^3\). The cervical lymph nodes are mainly involved in KFD, and the diagnosis is made from pathological findings of the resected nodes.

Only two cases of children with intra-abdominal lymph nodes involvement have been reported: a 14-year-old girl and a 15-year-old girl\(^4,5\). However, when making a differential diagnosis in patients with fever of unknown origin, KFD should be considered even if the surface lymph nodes are not palpable. We present here a case of KFD involving localized intra-abdominal lymph nodes because the diagnosis was difficult without a biopsy despite close evaluations aimed at identifying malignant lymphoma.

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CASE REPORT

A nine-year-old boy was admitted to the hospital because of persistent fever and intra-abdominal lymphadenopathy. The patient had been well until three weeks earlier, when fever developed. He consulted a clinic, and an antibiotic was prescribed. Ten days later, his fever went up to 39 °C and persisted. Although the antibiotic was changed, there was no clinical improvement. The patient consulted another hospital six days later and he was admitted because of a blood test showing a highly elevated transamisase. Cefotiam was administered intravenously, no improvement was observed. Because an abdominal CT scan showed swelling of the intra-abdominal lymph nodes he was referred to our hospital. He had had no night sweats or weight loss. There was no history of travel abroad or exposure to sick persons or animals, and there was no family history of allergic illness.

On examination, the patient appeared lethargic. His body height was 142cm and his body weight was 38.5kg. He had a temperature of 37.6 °C and his blood pressure was 130/80 mmHg. A physical examination was unremarkable; no petechiae, rash, or lymphadenopathy was found.

Fig. 1 A contrast-enhanced axial CT scan shows multiple enlarged mesenteric lymph nodes. (A)
An axial T2-weighted MR image shows homogeneous multiple lymphadenopathy. (B)
On a diffusion-weighted whole body image with background body signal supression, the enlarged lymph nodes showed slightly low signal intensity. (C)
Pathological finding of resected lymphnodes revealed abundant karyorrhectic debris and reactive histiocytes. (D)
His white blood cell count was 3,500/μL, with 49% neutrophils, 1% eosinophils, 36% lymphocytes, and 13% monocytes. His hemoglobin level was 12.1 g/dL, and his platelet count was 235,000 cells/μL. The serum levels of aspartate aminotransferase, alanine aminotransferase, alkaline phosphatase, lactic dehydrogenase, and C-reactive protein were 394 IU/L, 759 IU/L, 545 IU/L, 663 IU/L, and 0.39 mg/dL, respectively. His glucose, bilirubin, total protein, albumin, globulin, total cholesterol, blood urea nitrogen, creatinine, uric acid sodium, potassium, and chloride levels were normal. His ferritin was and soluble IL-2 receptor were 380 ng/ml and 875 U/ml. Natural killer cell activity was 15%. And β2 microglobulin in urine was 1648 μg/L. Bone marrow aspiration revealed normal findings except for some hemophagocytosis.

A CT scan of the abdomen disclosed enlarged mesenteric lymph nodes (Fig.1A). No organomegaly or ascites was seen. A chest CT scan was normal. Ga nuclear scanning revealed mild uptake in the midline region. Abdominal magnetic resonance imaging with gadolinium enhancement revealed multiple enlarged lymph nodes in the periaortic and supramesentric artery regions (Fig.1B). The regions revealed low signal intensity on short inversion time inversion-recovery imaging (Fig.1C). Although acetaminophen was given, the patient remained febrile. The mesenteric lymph nodes were biopsied by laparoscopy on seventh hospital day. Administration of diclofenac sodium, which was administered as an analgesic, relieved his fever and abdominal pain. However, when it was withdrawn on his 14th hospital day, his body temperature went up to 39°C, an eruption devel-

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Fig. 2 Clinical course
CTM: cefotiam  PSL: prednisolone
oped on his trunk and the bilateral conjunctivae became reddish. A histopathological examination of resected lymph nodes revealed abundant karyorrhectic debris and reactive histiocytes, and a diagnosis of KFD was made (Fig.1D). Prednisolone was administrated orally on the following day and his fever jell and the rash disappeared. He was discharged on his 29th hospital day. Thereafter prednisolone was tapered off. No symptoms and signs have recurred for a year (Fig.2).

DISCUSSION

The cause of KFD remains poorly understood. We found a moderate increase in serum soluble IL2 receptor (sIL2R). Since elevated levels of sIL2R have been reported in hematologic malignancies that express CD25 as well as in inflammatory diseases, especially in the active phase, measurement of sIL2R could not differentiate KFD from malignant lymphoma. The elevation of the sIL2R value could reflect activated T-cell responses, and moderate increases in ferritin and β2 microglobulin in urine might reflect hypercytokinemia, which causes leukocytopenia and hypertransaminasemia.

We attempted to diagnose the patient’s condition from imaging studies, but KFD has no specific radiological features. Kwon et al. observed lymph nodes with no specific signal intensity being usually affected, in their CT findings. In the magnetic resonance findings of our case, although the area of necrosis had a lower signal than non-necrotic areas on T2-weighted images, the intensity was homogeneous. Gallium-67 scintigraphy of the involved lymph nodes in this case showed uptake, as in the KFD patients described by Elshatifie et al.

Treatment for KFD is basically symptomatic treatment. Since KFD is benign and self-limiting, a “wait and see” policy is adequate. However, fever occasionally persists for months. Biopsy of the involved lymph nodes is warranted to make a definitive diagnosis, especially for differentiation from malignant lymphoma. In addition, fever subsides after excision in some patients. We administered prednisolone for our patients persistent fever after excluding malignant lymphoma.

In conclusion, we described a nine-year-old boy with persistent high fever and localized intra-abdominal lymphadenopathy which was diagnosed as KFD by biopsy. In this case, KFD could not be distinguished from malignant lymphoma by laboratory and diagnostic imaging findings. We recommend principal lymph node resection from diagnostic and therapeutic point of view. Laparoscopic procedures make biopsies safer and less invasive.

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