Kikuchi-Fujimoto disease mimicking tuberculous lymphadenitis or lymphoma on fluorine-18 fluorodeoxyglucose positron emission tomography/computed tomography

Noraini Abdul Rahim¹
Abdul Jalil Nordin²
Chiara Rusconi³
Cristina Gabutti³
Erika Ravelli⁴
Claudio Rossetti⁵

¹Diagnostic Imaging Department, Serdang Hospital, Selangor, ²Centre for Diagnostic Nuclear Imaging, University Putra Malaysia, Serdang, Malaysia, ³Department of Haematology, Ospedale Niguarda, Milan, ⁴Unit of Haematology, Valduce Hospital, Como, ⁵Department of Advance Technology (Nuclear Medicine), Ospedale Niguarda, Milan, Italy

Abstract: Kikuchi-Fujimoto disease, also known as histiocytic lymphadenitis is a rare and benign disease characterized by regional cervical lymphadenopathy with fever of unknown origin, affecting mainly young women. We describe a case of a young Asian female who was initially misdiagnosed as having tuberculous lymphadenitis because of multiple lymphadenopathies with 18-fluorodeoxyglucose uptake on positron emission tomography/computed tomography. Final histologic diagnosis of Kikuchi-Fujimoto disease was made after excisional biopsy of the affected lymph node. The cervical lymphadenopathy has reduced in size and lung lesions disappeared after 2 months without specific treatment.

Keywords: cervical lymphadenopathy, Kikuchi disease, PET/CT features

Case report
A 22-year-old female of Taiwanese descent presented to the Ospedale Niguarda Ca’ Granda, Milan, Italy with complaints of pain in the throat, associated fever, and sweats of 2 months’ duration. No significant past medical and surgical history was obtained. There was also no history of recent travels. Ten days later, she developed a 2 cm swelling in the right cervical region. Ultrasound of the neck showed multiple oval hypoechoic lesions with ill-defined margins seen in the right lateral cervical and parotid region. Ultrasound of the abdomen revealed no abnormality. The physical examination of other systems was normal. Subsequently, 1 month later, the patient presented again with fever, physical examination then revealed fullness of the nasopharynx and persistent lateral cervical lymphadenopathy. Patient was febrile. At this point, positron emission tomography/computed tomography (PET/CT) using fluorine-18 fluorodeoxyglucose (18F-FDG) tracer was performed and showed conglomerate FDG avid lymphadenopathies in the right cervical region and a well-defined solitary nodule in the upper lobe of the right lung with high FDG activity. Blood investigations revealed white blood count of 9,000/µL, neutrophil of 14.2 g/dL, platelet count of 404×10⁹/µL with normal liver and renal function tests. She was attended by the hematologist and received a course of oral antibiotics (Augmentin) which was changed to intramuscular Bactrim and Rocephin due to persistent temperature. Written informed consent was obtained from the patient before conducting the study. Ethical approval from the Ethics Committee of Ospedale Niguarda Ca’ Granda was not required for writing this case report.
The serum tests for toxoplasmosis and Epstein-Barr virus were negative. The serological titers for cytomegalovirus were positive. Purified protein derivative was positive. Serum immuno-stain markers correlated well with lymphoid hyperplasia. Biopsy of the roof of the nasopharyngeal lesion was done and histology was found compatible with lymphoid hyperplasia of the palatine tonsil.

Suspecting a lymphoproliferative disease, another 18F-FDG PET/CT was carried out which showed multiple small lesions in right lateral cervical and supraclavicular regions and two in the right pulmonary fields. A repeat ultrasound of the neck after 1 month showed multiple right lateral cervical nodes of increased dimensions compared to the previous examination. The underlying pathology was...
still uncertain. An excisional biopsy of the cervical node was done and histopathology examination reported past inversion of cellular architecture with immunochemistry positive to myeloperoxidase and partially to CD68r, tissue proliferative rate evaluation with MIB1 findings were compatible with Kikuchi-Fujimoto disease (KFD) with a high number of blast cells. Bone marrow aspiration and biopsy did not reveal any abnormality. Other causes of lymphadenopathies were also considered such as tuberculosis infection or other viral infections. Peripheral blood serology turned out positive for human herpes simplex virus IgM antibodies. Peripheral blood smear and autoimmunity tests were normal.

Considering the strong possibility of KFD, the patient did not receive any specific treatment. A repeat PET/CT performed after 2 months showed reduction in size and number of the cervical focal uptakes, and was negative in other areas including the right lung field. She remained well on review at the clinic 5 months later, and latest PET/CT after 9 months was found to be negative.

Discussion

KFD, also known as necrotizing histiocytic lymphadenitis, is a rare disease first described in 1972 by Fujimoto et al1 and Kikuchi.2 This disorder typically affects young women and shows a high prevalence among Asian people. It is a benign and self-limited disease, presenting with regional lymphadenopathies usually accompanied by constitutional symptoms like fever, frequently associated with upper respiratory tract symptoms. Less common symptoms are night sweats, malaise, fatigue, nausea, vomiting, and weight loss. The onset can be acute or sub-acute, usually developing during a period of 2–3 weeks, and at times lasting for 6–8 weeks. Etiology remains unclear. Many cases have reported associations with autoimmune disorders. Systemic lupus erythematous is often associated with it, thus in KFD patients close monitoring and follow-up to investigate for autoimmune diseases has been recommended. Some viruses are involved, in particular cytomegalovirus, Epstein-Barr virus, herpes simplex virus, and herpes zoster virus.

Because of the nature of the disease, it is challenging to diagnose KFD as it is frequently mistaken for malignant lymphoma. Final diagnosis is made by histological analysis of lymph node biopsy and no pathognomonic laboratory tests are available. It is characterized histopathologically by cortical and paracortical necrosis with lymphoreticular infiltration and absence of granulocytic infiltration.

There are no specific clinical or radiological features of KFD. The CT features of homogenously enhancing nodes without significant nodal necrosis make radiological differentiation from tuberculosis, metastases, or lymphoma impossible.

Lung involvement in KFD is unusual. Garcia-Zamalloa et al3 reported a case of pleural effusion and interstitial fibrosis complicating KFD. Fatal lung involvement has also been reported previously with cryptogenic organizing pneumonia and pulmonary hemorrhage.4,5 Migratory type of KFD was reported by Al Attia et al6 involving cervical, occipital, and jugulodigastric nodes. Recurrent KFD was sporadically mentioned in most case reports, but the actual rate is not known. It is also not known if KFD can transform into malignancy. In recent years, usefulness and usage of 18F-FDG PET in KFD has been reported. It has been shown that in KFD, lymph nodes have an uptake of FDG similar to aggressive lymphoma6–8 and other chronic granulomatous diseases like tuberculous lymphadenitis. This study can help clinicians to identify the severity of the disease through extension of lymph node involvement. Biopsy is mandatory, which is useful in assessing patient progress following treatment and confirming 18F-FDG PET/CT findings, especially during follow-up. The fact that KFD lesions demonstrate high 18F-FDG uptake on PET/CT, means that this can potentially be a future modality of choice for monitoring treatment response in KFD patients.

Conclusion

On diagnostic imaging including 18F-FDG PET/CT of the cervical lymphadenopathy in KFD, the findings can mimic lymphoma, metastases, or tuberculosis. In relatively young patients especially of Asian origin, KFD is a possible differential diagnosis and biopsy should be done to confirm the diagnosis.

Disclosure

The authors report no conflicts of interest in this work.

References


