Case Report / Olgu Sunumu

Langerhans cell histiocytosis: A rare cause of pathological rib fracture

Langerhans hücreli histiyositoz: Patolojik kaburga kırığının nadir bir nedeni

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ABSTRACT

Langerhans cell histiocytosis, formerly known as histiocytosis X, represents clonal proliferations of the antigen-presenting dendritic cells, which are normally found in many organs. It is a rare disease which tends to affect children and adolescents. In particular, adult-onset type is very rare. Herein, we present a female adult diagnosed with Langerhans cell histiocytosis of the rib without any systemic involvement which was successfully treated with surgery.

Keywords: Bone lesion; chest pain; Langerhans cell histiocytosis.

CASE REPORT

An otherwise healthy, 32-year-old female patient was admitted to the Department of Thoracic Surgery with right-side chest pain. The pain began one year earlier and was progressively getting worse, particularly over the last month. There was no history of trauma. Her medical history was non-specific and there was no family history of malignancy. Due to persistent pain, thoracic computed tomography (CT) showed a fracture line on the posterolateral area of the right 10th rib (Figure 1). A year-long progression of chest pain, pathological fracture line, and surrounding soft tissue enlargement indicated the possibility of a bone tumor. However, there were no abnormal hypermetabolic lesions on positron emission tomography (PET)-CT. A written informed consent was obtained from the patient. The 10th rib was removed by partial rib resection for diagnostic confirmation and treatment.
The pathology report was unexpectedly compatible with LCH (Figure 2). The rib resection material was 10×6×3.5 cm in total, consisting of the surrounding soft tissues and rib. The defined area was 3-cm away from the surgical margin. The lesion was composed of cells with large eosinophilic cytoplasm, vesicular nuclei, immunohistochemically positive staining for S-100, CD1a, and Langerin, as well as multinuclear giant cells stained with CD68 and diffuse eosinophils, which are clustered in places. Based on these findings, the patient was referred to the Department of Hematology.

During system evaluation, no dyspnea or tachypnea, polydipsia, and polyuria, lymphadenopathy, hepatosplenomegaly, weight loss, fever, gingival hypertrophy or ataxia was found. The body temperature, heart rate, blood pressure, and respiratory rate were all within the normal range. No clinical findings could be detected. Complete blood count and the blood
biochemical tests were within the normal range. Peripheral smear was examined, and no significant findings were found. A skeletal survey was negative for any other bone involvement. The endocrinological evaluation of the patient was made with blood tests and cranial magnetic resonance imaging (MRI), particularly in terms of diabetes insipidus and there was no pathological finding.

On radiological examination with whole-body skeletal survey and abdominal CT scan, there was no visceral lesion. The patient was thought to have unifocal involvement of LCH; therefore, no radiotherapy or chemotherapy was considered and was followed in the outpatient clinic. During follow-up for five months, the patient had no local recurrence or distant metastasis. She has remained asymptomatic during follow-up.

**DISCUSSION**

Langerhans cell histiocytosis includes a broad spectrum of clinical manifestations in children and adults, ranging from self-limited lesions to life-threatening disseminated disease. It may involve a single organ (single-system LCH), which may be a single site (unifocal) or involve multiple sites (multifocal) or it may involve multiple organs (multisystem LCH), affecting a limited number of organs or be disseminated.[4] In some case series, it is defined that bone lesions are predominantly observed in male patients, and other organ systems are involved in female patients.[4] The primary sites of bone involvement of LCH in adults, the ratios are as follows: mandible 30%, skull 21%, vertebrae 13%, pelvis 13%, extremities 17%, and ribs 6%.[5] It is most commonly seen in the head, but at least in the rib. Due to its similarity to radiological osteolytic lesions, LCH is difficult to differentiate from other bone tumors and benign lesions. The radiological findings of our patient were a fractured rib and coexistence of soft tissue swelling without osteolytic appearance. In such cases, it may come to mind whether these ratios are in patients with just solitary osseous lesion without extraosseous LCH or not, although there is no clear information about it. Also, lymphoma or lung carcinomas may be seen with LCH.[4] In our case, we observed no additional pathology or malignancy.

Considering the general characteristics of the disease, we consider that our patient with being a female adult and having a single bone lesion is one of the few cases in the literature.[5,9] Current data for the treatment of adults with LCH are limited to case reports and case series with no prospective clinical trials to inform therapeutic strategies.[1] Several numbers of approaches, including radiation, and various chemotherapy regimens, steroids, surgery or combinations of these treatment modalities, were used to treat LCH.[10] Surgical treatment includes commonly curettage of bone or excision of lymph nodes. Patients with a solitary osseous lesion have the best prognosis compared to those with LCH involvement of other systems.[4] In our case, partial resection of the rib was performed, as the patient had persistent pain for months and a pathological fracture line surrounding soft tissue enlargement. After examination, no additional involvement site was detected in our patient and diagnosed with a single lesion-LCH.

In conclusion, surgical excision is kept in mind as an effective treatment for Langerhans cell histiocytosis, particularly in patients with solitary bone lesions.

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