A Langerhans cell histiocytosis in its rare Letterer-Siwe form in a child mimicking osteoarthritis: A case report and literature review

Bouali Mustapha, H Mezhoud, H Soltani and I Fergoug
Oran University Hospital, Algeria

Langerhans cell histiocytosis (LCH) is a group of three rare diseases (annual incidence less than 4/1000,000 H) with increasing aggressiveness respectively: Eosinophilic granuloma (unifocal solitary osteolytic lesion), Hand-Schuller-Christian disease (multifocal lesion) and Letterer-Siwe disease (the most aggressive, disseminated disease with systemic manifestations). The bone is most commonly involved with a predilection for the skull. Some cases of temporal bone localizations have been described in the literature. We report a case of an 18-month child presenting a LCH of the temporal bone misdiagnosed at the beginning as osteoarthritis. In fact, the child presented a retro-auricular swelling with inflammatory characters, fever and otorrhea. The exploration before surgery revealed besides the extensive temporal lesion, hematologic perturbations, a splenomegaly and a hepatomegaly. The histopathology confirmed the diagnosis of a LCH. We describe the diagnosis challenges, the surgical difficulties and the therapeutic response of the child.

Notes: