**Langerhans Cell Histiocytosis with Multiple Fluid–fluid Levels in the Parietal Bone**

Kazuki Matsushita*, Taro Shimono, and Yukio Miki

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**Introduction**

Langerhans cell histiocytosis (LCH) is the most common histiocytic neoplasm involving the calvarial bone in children. LCH characteristically shows almost homogeneous signal intensity on gadolinium-enhanced MR imaging. However, LCH manifesting as an osteolytic cystic mass with multiple fluid–fluid levels (FFLs) is rare. This unique imaging appearance has rarely been reported. Herein, we report a case of solitary LCH in the skull with FFLs.

**Case Report**

A 10-year-old girl presented with a 3-week history of an occipital mass with persistent swelling and swelling and tenderness for 2 weeks after bruising her head. No other abnormalities were found on clinical examination. Skull radiographs demonstrated a single osteolytic lesion without sclerosis in the left parietal bone. Computed tomography demonstrated a mass that extended beyond the cortex of the inner and outer tables (Fig. 1). A beveled edge appearance was demonstrated due to asymmetrical involvement of the inner and outer tables. T2-weighted images showed multiple FFLs (Fig. 2a), whereas T1-weighted images showed a broad signal intensity change in the adjacent bone marrow (Fig. 2b). Post-contrast T1-weighted images showed enhancement of the solid part of the lesion and adjacent pericranial soft tissue, dura mater, and bone marrow (Fig. 2c).

During surgery, an extradural encapsulated cystic mass containing hematoma was found beneath the periosteum. Pathologic examination of hematoxylin–eosin stained sections revealed proliferation of plasma cells, lymphocytes, and histiocytic cells. The histiocytic cells were confirmed to be Langerhans cells by their positive immunohistochemical staining for CD1a, langerin, and S100 protein. Thus, the diagnosis was LCH.

**Discussion**

Langerhans cell histiocytosis characteristically shows almost homogeneous enhancement on MR imaging, whereas in our patient it was manifested as a bony mass lesion containing small cystic portions with FFLs. Nabavizadeh et al. reported 11 pediatric patients with skull masses containing FFLs. The underlying pathology was LCH in four children, aneurysmal bone cysts (ABCs) in three, cephalohematoma in three, and metastatic neuroblastoma in one. Though FFLs are famous characteristics of ABCs, the beveled edge appearance in this case cannot be seen in ABCs. These FFLs in LCH represented hemorrhage, which were caused by trauma or developed spontaneously. Therefore, the top differential diagnosis for cystic skull lesions with FFLs in children is LCH.

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Department of Diagnostic and Interventional Radiology, Osaka City University Graduate School of Medicine, 1-4-3 Asahi-machi, Abeno-ku, Osaka, Osaka 545-8585, Japan

*Corresponding author, Phone: +81-6-6645-3831, Fax: +81-6-6646-6655, E-mail: m2075787@med.osaka-cu.ac.jp

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**Fig. 1** Computed tomography image showing a mass (arrow) extending through and beyond the cortex of the inner and outer tables in the left parietal bone. A beveled edge appearance was demonstrated due to asymmetrical involvement of the inner and outer tables.
Our patient also showed T1-weighted low-intensity changes in adjacent bone marrow that resolved after resection of the lesion. Stull et al.\textsuperscript{3} have reported that signal-intensity alterations within surrounding marrow suggest either reactive edema or inflammation. Okamoto et al.\textsuperscript{4} reported similar changes of calvarial LCH. Thus, these reactive changes on MR imaging may be characteristic of LCH, and unlikely imaging findings for malignant tumors accompanied with FFLs.

We consider awareness of these unique presentations of LCH on MR imaging is important to know.

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Conflicts of Interest

The authors declare that they have no conflicts of interest.

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