Temporal Brain Hydatid Cyst in an Eight-Year-Old Child: A Rare Case Report

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Abstract: Hydatidosis is a parasitic disease caused by Echinococcus granulosus, which affects children in many different parts of the world. It commonly affects the lungs and the liver of the children. Brain hydatidosis is an extremely rare clinical condition in the pediatric population, presenting with non-specific symptoms and signs. The diagnosis of intracranial hydatid cysts can be established by brain magnetic resonance imaging and histopathological examination of the specimen. Here, we report an 8-year-old female child diagnosed with a temporal brain hydatid cyst. Brain magnetic resonance imaging showed a thin-walled cystic lesion located in the right temporal lobe with significant mass effect and midline shift, with no abnormal wall or solid enhancement, and no surrounding edema. The diagnosis of brain temporal hydatid cyst was made radiologically. The patient was operated on and the cyst was completely removed without rupture. The removed cyst was sent for histopathological examination; the histological sections showed a laminated acellular cyst wall with a nucleated germinal layer and no protoscolices, and the diagnosis of temporal brain hydatid cyst was confirmed. The patient had a smooth postoperative course, started albendazole therapy, and was discharged with improvement.

Keywords: temporal brain hydatid cyst, Echinococcus granulosus, histopathology, cyst wall, albendazole

Case Presentation

An 8-year-old female Ethiopian child presented to the hospital with a 4-month history of global headache, which worsened progressively. She also had projectile non-bilious vomiting two or three times a day and blurring of her vision for a duration of 1 month. Otherwise, she had no abnormal body movements or gait disturbances.

Physical examination revealed intact cranial nerves and no sensory or motor deficits. Chest X-ray and abdominal ultrasound were normal. Magnetic resonance imaging (MRI) of the brain showed a cystic signal intensity similar to that of cerebrospinal fluid (CSF), without ring enhancement. The axial T1-weighted brain MRI showed a well-defined hypointense lesion in the right temporal lobe with significant mass effect and midline shift (Figure 1). Coronal T2-weighted and axial fluid-attenuated inversion recovery (FLAIR) images of the brain showed a well-defined T2 hyperintense lesion that was totally suppressed on the FLAIR sequence, with no sign of perilesional edema (Figure 2).

The sagittal and axial T1-weighted MRI of the brain after gadolinium administration showed a well-defined cystic mass with no enhancement on the cyst wall, and there was no mass-like or mural nodular enhancement (Figure 3).

Based on the presentation and brain MRI findings, the diagnosis of giant intracranial hydatid cyst was made. Right temporoparietal craniotomy was carried out and a huge cyst, measuring 15×13×12 cm, was removed with an intact capsule, with maximum care taken to avoid rupture and spillage (Figure 4).

The removed cyst was sent for histopathological examination. The gross section showed a 7×6×4 cm whitish cystectomy specimen with a smooth outer surface. On cut section, it was found to be a unilocular cyst filled with...
clear fluid and with a rough inner surface. The wall thickness was 1–2 mm. Microscopically, the histological sections showed a laminated acellular cyst wall with a nucleated germinal layer. No protoscolices were seen (Figure 5).

Postoperatively, the patient was stable. Albendazole 15 mg/kg was initiated and continued for 4 weeks. The patient showed marked improvement with resolution of her symptoms.

**Discussion**

Hydatidosis is a parasitic disease caused by *Echinococcus granulosus*, which affects children in many parts of the world. Brain involvement with cystic hydatid disease occurs in 1–2% of all *Echinococcus granulosus* infections. Intracranial hydatid cysts constitute one of the major etiologies of intracranial space-occupying mass lesions in endemic regions, especially in children and adolescents. The clinical presentation of brain hydatidosis depends on the size of the cyst and
Figure 3  T1-weighted MRI of the brain after gadolinium administration showing a well-defined cystic mass with no enhancement on the cyst wall.

Figure 4  The removed giant intracranial hydatid cyst after the operation.

Figure 5  Histopathology microscopic section showing a laminated acellular cyst wall with nucleated germinal layer.
its location. The most commonly reported symptoms are headache, vomiting, body weakness, abnormal body movements, behavioral alterations, and skull deformity.\textsuperscript{3} Brain hydatid cysts may grow up 1–5 cm in a year and the presenting clinical signs are often related to elevated intracranial pressure and focal neurological deficit.\textsuperscript{3,4}

Imaging is crucial to establish the diagnosis and guide the treatment of children with intracranial hydatid cyst. Brain CT and MRI showed a well-defined spherical homogeneous cystic lesion with a thin wall and smooth margins, and imaging characteristics of the cystic component similar to that of CSF. On MRI, the cyst wall usually has low signal intensity on both T1-weighted and T2-weighted images. Calcification of the wall is uncommon, reported in <1% of all cases.\textsuperscript{5,7,8} The diagnosis of brain hydatidosis is confirmed by histopathological examination of the specimen. The cyst wall has three layers. The inner germinal layer has a thickness of 10–25 µm. The cuticular membrane is followed by a laminated membrane and then fibrovascular tissue.\textsuperscript{6}

The primary treatment of brain hydatid cysts is surgery and the main goal of the neurosurgical intervention is to excise the cyst without rupture to prevent recurrence and anaphylactic reaction.\textsuperscript{9} Postoperatively, albendazole therapy is recommended to be given for 1–3 months.\textsuperscript{10}

Our case report showed a rare presentation in an 8-year-old child with a giant intracranial hydatid cyst. Brain hydatidosis should be considered as a differential diagnosis in cystic cerebral mass lesions in the pediatric age group. She was operated on and the cyst was removed completely without rupture. The histopathological examination showed a 1–2 mm thick unilocular and laminated acellular cyst wall with a nucleated germinal layer. Postoperatively, she had no complications and was discharged home with albendazole therapy.

**Conclusion**

Although hydatidosis remains a significant issue worldwide, intracranial hydatid cysts are rare in clinical practice. Diagnosis is based upon history, imaging, and histopathological analysis. Neurosurgical excision is the mainstay of treatment for intracranial hydatidosis. Early diagnosis and timely interventions are important to avoid acute life-threatening complications and long-term sequelae.

**Ethical Clearance**

Written informed consent for publication of their details, including the intraoperative images and histopathology, was obtained from the parents. In addition, ethical clearance was obtained from Tikur Anbessa Hospital and Addis Ababa University institutional review board.

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**Disclosure**

The authors report no conflicts of interest in relation to this work.

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