Primary Hydatid Cyst of the Brain in a Child: A Case Report

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Summary

Background: Primary intracranial hydatid cyst is a rare location of human echinococcosis whose spontaneous, traumatic or even iatrogenic rupture, as in case of misdiagnosis, may cause anaphylactic reactions and dissemination.

Case Report: We discuss the management of a 9-year-old boy who was admitted to our Emergency Department with an intracranial hypertension syndrome. Head CT scan and brain MRI showed a huge intra-axial right temporo-parieto-occipital cyst with a marginal calcification, associated with left ventricular uncompensated hydrocephalus. DTI showed displacement of the ipsilateral corticospinal tract, whereas MR spectroscopy showed absence of normal brain metabolites and presence of succinate and lactate within the cyst. A diagnosis of hydatid cyst was then presumed on the basis of the neuroradiological findings. Empiric chemotherapy with albendazole was instituted and surgical en bloc removal of the cyst was obtained, allowing the patient to recover without complications. Diagnosis of brain echinococcosis was confirmed by laboratory tests.

Conclusions: HE is still an endemic manifestation in some rural areas of the world, and it should be included in the differential diagnosis of children living in or coming from an endemic country who present with an intracerebral cyst. Early diagnosis and complete surgical removal of the intact cyst are the main factors that determine a favourable outcome.

MeSH Keywords: Brain Diseases • Child • Echinococcosis

Abbreviations: HE – human echinococcosis; CT – computed tomography; MRI – magnetic resonance imaging; CSF – cerebrospinal fluid; DTI – diffusion-tensor imaging

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Spontaneous or traumatic (including surgical) cyst rupture may cause anaphylactic reactions and dissemination [3,4]. We present the case of a giant primary intracerebral hydatid cyst in a child, discussing its surgical management and analyzing the recent literature on the topic.

**Case Report**

We examined a 9-year-old boy who recently immigrated from a rural area of Albania, who complained of severe long-standing headache and diplopia. He was alert and conscious but asthenic, without sensorimotor deficits. Fundoscopy revealed left papilledema, while routine blood tests, chest X-rays, and abdominal ultrasound were unremarkable. Cardiac ultrasound revealed a patent Botallo duct with left-to-right shunt. Head computed tomography (CT) scan showed a huge intra-axial right temporo-parieto-occipital hypodense cyst with marginal calcification. Brain magnetic resonance imaging (MRI) confirmed the cyst to be isointense with cerebrospinal fluid (CSF) and causing marked mass effect, contralateral displacement of the midline structures, and left ventricular uncompensated hydrocephalus (Figure 1). Diffusion-tensor imaging (DTI) showed displacement of the ipsilateral corticospinal tract, whereas MR spectroscopy showed absence of normal brain metabolites and presence of succinate and lactate within the cyst (Figure 2). A diagnostic hypothesis of hydatid cyst was made on the basis of the neuroradiological findings and empiric chemotherapy with albendazole was instituted. Further history taking revealed that he routinely used to drink raw non-pasteurised milk. The patient underwent surgery via a large occipital craniotomy; a thin cortical layer covered the cyst. Gross total removal was obtained via Dowling’s technique [5,6], without rupture of the wall and/or ventricular opening. The patient recovered without complications and post-operative CT and MRI confirmed complete removal of the lesion. The patient was discharged home after suture removal. Cystic fluid was analyzed and protoscoleces were stained (Figure 3). Two months after the operation, persistent Botallo duct was successfully corrected through an endovascular approach and therapy with albendazole was discontinued eight months later.

**Discussion**

*Echinococcus granulosus* and *Echinococcus multilocularis* are small tapeworms that live in the bowel of canines (definitive host); their eggs are shed in the feces of dogs and can be ingested by cattle, sheep, and sometimes humans (intermediate host) [1]. After digestion of the capsule,
Figure 2. (A) Axial diffusion-weighted image and (B), corresponding apparent diffusion coefficient (ADC) map reveal water-like diffusion within the cyst. Placement of an MR spectroscopy voxel is also shown on A. (C) Color-coded fractional anisotropy image derived from a diffusion-tensor imaging (DTI) study at 16 directions shows that the corticospinal tract (CST) is severely distorted and displaced anteriorly (arrows; compare with normal contralateral CST, arrowheads). Fiber tractography image (E), however, confirms that the CST is only displaced, but not infiltrated, by the adjacent mass. (D) MR spectroscopy obtained with a single-voxel point-resolved (PRESS) technique at 144 ms echo-time shows absence of normal brain metabolites within the cyst; a huge peak at 2.4 ppm is consistent with succinate, whereas an inverted double peak at 1.3–1.4 ppm is consistent with lactate.

Figure 3. Protoscoleces from the hydatid cyst fluid, stained with Dobell (40× magnification). Note the hooklets (black arrow).
embryos penetrate the intestinal mucosa and spread through the portal circulation: most are trapped in the liver and lung, developing hydatid cysts in those locations. The brain is a rare location, accounting for only 1–2% of all intracranial space-occupying lesions even in endemic areas [2]. Associated extracranial cysts are common, thus requiring a thorough radiological evaluation of the patient (including chest X-rays, abdominal ultrasound and, especially in children, cardiac ultrasound).

Primary cerebral HE is exceedingly rare; only few reports from endemic areas such as Turkey and India are available [7,8]. To date, only one case report and one clinical series have been described by Lunardi et al. and Sardi et al. of Italian patients operated on for hydatid brain lesions [9,10]. Brain location requires peculiar anatomic conditions that allow the germ to shunt the hepatic and pulmonary filters; the present patient had a persistent Botallo duct [11]. Most reported cases have described solitary, intraparenchymal cysts mainly located along the terminal branches of the middle cerebral artery [12]. Especially in children, cerebral hydatid cysts can grow to an enormous size because of the elastic structure of bone and the relatively higher compressibility of the neural tissue [13]. Multiple cysts can be due either to embolization of multiple larvae or to traumatic, iatrogenic, or even spontaneous cyst rupture [14,15]. Clinical features are not specific, depending on size and location: headache and signs of raised intracranial pressure are mostly reported, while focal neurological deficits and seizures are less frequent. Ersahin et al. reported a growth rate of about 1 cm per month over a 6-month period [16]. Serologic tests are of little clinical value in primary cerebral disease [13,17].

MRI and CT scans are helpful in suggesting the diagnosis and planning a proper treatment, although no pathognomonic sign can be detected [18]. The cyst characteristically presents as a solitary, homogeneous, spherical lesion with well-defined borders, no contrast enhancement, and no perilesional edema, filled with CSF-isodense fluid. Other supratentorial cystic lesions such as arachnoid cysts, cystic tumors, abscess, and porencephalic cysts should always be considered in the differential diagnosis [19]. Advanced imaging techniques, such as DTI or MR spectroscopy, can be helpful but not conclusive.

Surgery is the elective treatment, the goal being removal of the entire cyst without rupture, thereby preventing parasitic spread with further neurological deficits and the rare but existing anaphylactic reaction [3,20]. This can be achieved following some simple, clear steps: (i) positioning the patient’s head so that the cyst is in a dependent position; (ii) large craniotomy; (iii) adequate corticotomy with accurate cortical dissection; (iv) careful Valsalva manoeuvre; and (v) use of continuous warm saline irrigation between the cyst and the surrounding brain parenchyma to help delivery (Dowling’s technique) [5,6]. Other techniques have been proposed, such as fluid aspiration, but they appear to be dangerous with a high probability of cyst rupture and associated anaphylaxis and/or parasite spread to distant sites [4]. Intraoperative cyst rupture is a possible event associated with a potentially fatal anaphylactic reaction and a high rate of cyst recurrence [21, 22]. On the other hand, seizures, subdural effusion, porencephalic cyst, hemorrhage, pneumocephalus, hydrocephalus, and focal neurological deficits are the main reported postoperative complications that sometimes need further intervention [23]. Medical therapy is also important. Albendazole is currently the elective medication; it is effective in sterilizing the cyst, decreasing the risk of anaphylaxis, and reducing the recurrence rate [24,25]. Corticosteroids may help control perilesional edema, while anticonvulsants are used prophylactically [26].

Conclusions

HE is still an endemic manifestation in some rural areas of the world, and it should be included in the differential diagnosis of children living in or coming from an endemic country who present with an intracerebral cyst. Early diagnosis and complete surgical removal of the intact cyst are the main factors that determine a favourable outcome. When a primary intracranial hydatid cyst is strongly suspected, the treatment of choice is surgery. The best surgical technique is en bloc removal of the cyst. Cyst puncture should be avoided for the risk of anaphylactic reaction and parasite spread to distant sites.

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