Incidentally Detected Cardiac and Hepatic Hydatid Cyst after Sudden Onset Facial Paralysis: A Case Report

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Abstract
The simultaneous occurrence of interventricular cardiac septum and hepatic localization for cystic echinococcosis is a very rare form of presentation and it may lead to various complications. We present the case of a 24-year-old man who was admitted to the hospital for evaluation of abrupt onset facial paralysis and incidentally diagnosed with a large hepatic hydatid cyst and a cardiac one located in the interventricular septum. The patient did not have any other symptoms associated with hydatid disease before this event. In this case, we surgically removed both hydatid cysts at the same time, to prevent complications and to establish a better outcome for the patient.

Keywords: Cardiac cyst; Hepatic cyst; Hydatidosis; Facial paralysis

1. Introduction
A flat tapeworm, Echinococcus granulosus, causes hydatid disease in humans, a parasitic disease that occurs in some endemic regions of the world. The disease is caused by the larval stage of E. granulosus. The life cycle alternates between carnivores and herbivores. The life cycle of this parasite involves dogs as definitive hosts. Eggs are passed in the faeces and eaten by the intermediate hosts, such as sheep, and the larvae encyst in the liver, lungs, and other organs. The natural cycle completes when a dog eats the sheep viscera and ingests the larvae cyst. The protoscolices attach to the small intestinal wall and the worms begin to form proglottids which will be containing the
eggs. The eggs pass in the faeces. Human beings act as an intermediate accidental host, and an end point in the parasite’s life cycle. Human infestations develop after accidental ingestion of excreted eggs in animal faeces. Consumption of unwashed and raw contaminated vegetables is the source of the infestation. The embryo develops from the egg and migrates via the intestinal mucosa into the circulation. Any organ can be involved by the cyst formation, most frequently the liver (65%) and lungs (25%) [1, 2]. Cardiac hydatid cysts are found in fewer than 2% of cases of hydatidosis [3]. The most common cardiac locations are the left ventricular wall (60%) followed by the right ventricle (10%), pericardium (7%), left atrium (6-8%), right atrium (4%), and the interventricular septum (4%) [4]. In 50% of such cardiac cases, there is a multiple organ inclusion [4]. Cardiac hydatid cysts grow slowly, the disease may be asymptomatic for a long period of time, is usually detected incidentally on echocardiography taken for other reasons. A definitive diagnosis is made by use of computed tomography (CT), magnetic resonance imaging (MRI), in addition to serologic tests [5]. The disease is treated by a combination of surgery and medical therapy with Albendazole.

2. Case Report

A 24-year-old male was referred to our hospital from a neurology clinic where he had been evaluated for chief complaint of abrupt onset facial paralysis and mild headaches. Physical examination in our clinic, one week after the onset of the symptoms, was unremarkable, with exception of the neurological examination that showed a mild facial paralysis. Past medical and surgical history were unremarkable. Social history revealed that the patient has been in close contact with animals such as sheep and canines by the nature of his occupation. His initial vital signs were normal, the temperature was normal. Laboratory test findings only showed neutrophilia (WBC count at 14.710/μL, neutrophil count at 8.470/μL) and ALT of 55 u/L (normal range 10-50 u/L). The electrocardiogram (EKG) showed sinus rhythm at 80/min, QRS axis at +45°, negative T waves in DIII lead, no other ST-T abnormality. Cerebral tomography done at the referring hospital a week prior to admission revealed subcortical, thalamic, mesencephalic and hypothalamic small ischemic lesions, some of them with hemorrhagic transformation. Transthoracic echocardiography (TTE) showed a cystic structure measuring 23/19 mm located at the junction between the apical and the middle third of the interventricular septum toward its anterior part (Figure 1).

At this level, a mobile mass measuring approximately 3/1,5 cm was attached to the hyperechogenic wall of the cystic structure, highly suggestive for thrombus. A new head CT was obtained at our clinic. Hemorrhagic cerebral lesions were stable as compared to the prior head CT. The neurological evaluation raised the suspicion of facial paralysis frigore, and in the context of the mobile mass attached to the cystic cardiac wall, it is considered appropriate to start anticoagulant treatment. Continuous infusion of unfractionated heparin was started. A full-body CT-scan was performed and confirmed the presence of the cardiac cyst (Figure 2). Consequently, a second larger cystic formation was discovered located in the hepatic parenchyma, measuring 62/47 mm (Figure 3). Serologic tests for Echinococcus granulosus were sent as clinical suspicion was very high for hydatid disease. Empiric Albendazole (400 mg twice daily) treatment was thus initiated. Serologically, specific Echinococcus granulosus (E. granulosus)
and *Echinococcus multilocularis* (*E. multilocularis*) antibodies were investigated by commercial Enzyme-Linked Immunosorbent Assay (ELISA) test. *E. multilocularis* antibodies were positive by ELISA. Hydatid cyst antibodies had positive 1/320 titer in the Hemagglutination Inhibition Test (HAI). As expected, ELISA were highly positive: 5.37 (normal range <0.9) for echinococcus antibodies. With Albendazole treatment, leukocytosis resolved. Overall status remained very good. Since the patient was 24 years old and did not have any risk factors for coronary atherosclerosis, further investigations were not performed.

Surgery was scheduled after cerebral CT examination showed stabilization of the hemorrhagic lesions in order to minimize the cerebral hemorrhage risk during CPB. We decided to remove first the cardiac hydatid cyst. The patient underwent median sternotomy and was placed on standard aorto-bicaval cardiopulmonary bypass. After aorta was cross-clamped, for cardiac arrest we used cold potassium-enriched blood cardioplegic solution. The left atrium was vented through the right upper pulmonary vein. The pericardial cavity was wrapped with sponges moistened with 20% hypertonic saline solution and povidone iodine 10% solution (larvicidal solution). The incision to excise the cyst was made at the apical left lateral wall ventricle parallel to the left anterior descending coronary artery, to avoid damaging that vessel. A 2-3 cm well-incapsulated hydatid cyst was found in the proximal third of the interventricular septum (Figure 4). We found the cyst not to be covered by the thrombus. A syringe with 50 ml of larvicidal solution was introduced into the cystic sac. After 5 minutes the cavity was aspirated with the same syringe and the cyst cavity was opened and the protoscoleces (fertile cysts) derived from the germinal layer was carefully removed completely. The edge of the cyst was excised and the cavity in ventricular septum was washed with a larvicidal solution and left open. The left ventriculotomy was closed with 3-0 polypropylene continuous sutures over Teflon felt. At the end, the pericardial cavity was washed with a larvicidal solution. The patient was weaned from cardiopulmonary bypass, intraoperative TEE examination showed normal biventricular function without any traces of intraseptal cardiac hydatid cysts. In the same surgical procedure we also removed the hepatic cyst. Before closing the chest, laparotomy was performed with exposure of the surface of encapsulated hepatic hydatid cyst. In similar fashion, the hepatic cyst was wrapped with sponges moistened with larvicidal solution. The cyst was punctured with a wide needle and 100 cm$^3$ of the cystic content was aspirated, then the cyst content was sterilized by mixing the remaining contents with the larvicide solution. After a few minutes, the cyst was completely aspirated and opened. The cyst content, endocyst and remaining daughter cysts should be enucleated from the pericyst with a sharp spoon and the residual cavity was washed again with a larvicidal solution (Figure 5). The patient had an uneventful postoperative course, with a minimal intensive care unit stay and was discharged in the 7th day postoperatively with 400 mg albendazole twice a day for the next six weeks postoperatively. One year follow-up showed no pathological findings.
**Figure 1:** Echocardiographic aspect of the hydatid cyst located in the interventricular septum.

**Figure 2:** CT showing a cystic lesion on the interventricular septum.

**Figure 3:** CT showing a cystic lesion of the liver.
Figure 4: Intraoperative view of the cardiac cyst.

Figure 5: Intraoperative view of the remaining cavity after the inactivation of the cardiac cyst.

3. Discussion

Hydatidosis is a parasitic infection caused by *Echinococcus granulosus*. After infestation, the embryo usually reaches the myocardium via the coronary circulation from the left side of the heart and reaches the liver via intestinal mucosa into the portal circulation. Enlargement in the cyst volume requires a period of one to five years [6] and can compromise adjacent structures, thus causing symptoms. During this time it may calcify and die, or grow in situ. Due to slow disease progression, patients suffering from hepatic and cardiac echinococcosis may be initially asymptomatic for a long time. The clinical manifestation may vary according to the location, size and age of the cysts and the extent of calcification. When symptomatic, patients generally present with palpitation, dyspnea, atypical angina pectoris, cough, haemoptysis, compression with acute coronary syndrome, valve obstruction and biventricular systolic dysfunction, cardiac syncope, arrhythmias and conduction disorders, acute pericarditis, or cyst
rupture with cardiac tamponade [7], anaphylactic reaction with profound circulatory collapse and cerebral or peripheral arterial embolism [8], pulmonary embolism with pulmonary hypertension [7] or even sudden death [9]. The most frequently involved cardiac region is the left ventricle while the right ventricle and the interventricular septum are the less affected regions, respectively [10].

Our patient was asymptomatic for years and did not have any complaints associated with hydatid disease before the sudden occurrence of the facial paralysis accident with which the cardiac cyst has been diagnosed. The first problem was diagnosed the etiology of facial paralysis. Repeat cerebral CT revealed small ischemic lesions, stationary in evolution. TTE showed a cystic structure on the left side of the interventricular septum with a mobile mass attached, highly suggestive for thrombus. We considered the small cerebral ischemic lesions detected on cerebral CT to be the result of the thrombus embolising from the mobile mass attached to the wall of the cardiac hydatid cyst. Since facial paralysis was not accompanied by any other neurological sign and the ischemic lesions were small and disseminated, we suspected that it was rather a paralysis a frigore than the result of ischemic stroke. The second diagnostic problem after ultrasound identification of a cardiac tumor mass was the possible confusion with the simultaneous development of left ventricular myxoma or papillary fibroelastomas with concomitant hepatic hydatid cyst [11].

However, the literature describes left ventricular myxoma only in case of recurrence of the disease [12, 13]. Specific diagnosis of hydatid cyst in general is based on clinical presentation, serology tests and non-invasive imaging techniques such as TEE, CT and MRI. The ELISA is one of the most specific serologic tests that can be used and a positive result for Echinococcus antibodies confirms the diagnosis [14]. Serologic tests can be false-negative in 10% to 20% of patients with hepatic hydatid cysts and 50% with cardiac cysts; this is most likely linked with an insufficient immune response [15]. Echocardiography remains the most efficient method in the diagnosis, which provides important findings: size and number of cysts, cyst locations and relationship with adjacent structures [1]. In our case, the positive serological tests confirmed the presence of hydatidosis. Due to the high risk of the spontaneous rupture of cysts or systemic embolisms, the treatment of choice, even in asymptomatic cases, is the complete removal of cysts with early cardiac surgery under cardiopulmonary bypass. From the current literature, an estimation of prognosis for patients that received non-surgical treatment is not possible. For cardiac location, there is a case report of surgical treatment on beating heart when the cyst is located on the epicardium [16]. In our case, both CT and echocardiographic findings were indicating intraseptal involvement so the cardiopulmonary bypass by conventional technique was used. When a patient presents with hydatid cysts in cardiac and alternate location, it is possible to elect to combine or not the cardiac surgery with laparotomy.

The general principle is to give priority to the cysts before complications occur. In our case, the emergency was to remove the cardiac cyst that was responsible for stroke. The incision to excise the intraseptal cyst must offer direct access and can be made from the atrium, ventricle or apex cordis. In our case, we prefer an incision in the left apical
lateral ventricle. However, there are also reports with sudden cardiac death after resection of the cardiac hydatid cysts [17]. The contents of the cyst must be entirely aspirated and the germinative membrane removed. Although many larvicidal agents have been introduced in order to prevent dissemination and recurrence is widely used, none is 100% effective and safe for intraoperative use. We used a relatively nontoxic protoscolicidal agent, a combination between 20% hypertonic saline solution and 10% povidone iodine solution. This combination destroys protoscolices by creating a significant osmotic gradient [18]. We have also used this solution to isolated surrounding areas to prevent the risk of development of secondary cysts as a result of leakage from the primary cyst during the surgical procedure and to wash out the remaining cavity.

Along with the surgical treatment, currently the use of the synthetics benzimidazole-derivative anthelmintic has been shown to be effective in the treatment of hydatid disease and to prevent recurrence of cysts after operation [19]. Medical treatment should be started a few weeks prior to surgery, yet for cardiac cysts that therapy may lead to a higher risk of cyst wall destruction and rupture [20]. Surgical resection should be followed by therapy with albendazole (10–15mg/kg body weight per day) or mebendazole (40–50 mg/kg body weight per day). In our case, the patient was treated preoperatively for 15 days with albendazole 400 mg twice per day and was discharged with the same dose for the next six weeks postoperatively. About 10% of all hydatid cysts tend to recur after surgery, but this rate may decrease with albendazole therapy. However, albendazole therapy is contraindicated for large cysts with the risk of rupture and also for inactive or calcified cysts [21].

4. Conclusions
Simultaneous cardiac and hepatic hydatid cysts are rare and often asymptomatic in their early stages. Clinical suspicion is important in reaching an early diagnosis. In patients with cardiac masses the possibility of cardiac hydatid disease should be taken into consideration, even outside an endemic area. Serology tests, TEE, CT and MRI are useful for diagnosis and establishing the location of hydatid cysts. Due to the high risk of their associated complications, hydatid cysts should be removed surgically, even in asymptomatic patients.

Conflict of Interest
The authors declare that they have no competing interests.

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References


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