A Large Intracardiac Hydatid Cyst with Concomitant Cervical and Hepatic Involvement: A Case Report

Maryam Faramarzpour¹, Sirous Jafari², Mehrzad Rahmanian¹, Akram Sardari¹, and Farnoosh Larti¹

¹Tehran University of Medical Sciences
²Imam-Khomeini Medical Center

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Abstract

Cardiac hydatidosis is a relatively rare complication of echinococcosis, with a potentially life-threatening condition. Here, we reported a large interventricular septal hydatid cyst with bulging in the left ventricle accompanied by a huge cervical lump with recurrent hepatic cysts that underwent cardiac surgery to excise the cyst uneventfully.

Title page

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Author’s names:

1. Maryam Faramarzpour¹, MD
2. Sirous Jafari², MD
3. Mehrzad Rahmanian³, MD
4. Akram Sardari¹, MD
5. Farnoosh Larti*¹, MD

*Address for correspondence: Farnoosh Larti, MD. Postal address: Imam Khomeini Hospital Complex, Tohid Square, Tehran, Iran Postal Code: 1419733141 Tel: 00989118319061, Fax number: 00982166939537 Email: Farnooshlarti@gmail.com

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Key Clinical Message

Cardiac hydatidosis is a relatively rare complication of echinococcosis. Understanding the atypical manifestations, potential associated risk factors, and epidemiology leads to optimal and timely management.

Abstract

Cardiac hydatidosis is a relatively rare complication of echinococcosis, with a potentially life-threatening condition. Here, we reported a large interventricular septal hydatid cyst with bulging in the left ventricle accompanied by a huge cervical lump with recurrent hepatic cysts that underwent cardiac surgery to excise the cyst uneventfully.

Introduction

A 68-year-old woman with a history of hepatic hydatidectomy two years ago complained of a large right-sided cervical mass. In ultrasound imaging, a huge (45 mm × 75 mm) mass was detected in the base of the right side of the neck. Moreover, multiple cystic lesions involving the two lobes of the liver were found. Despite the absence of cardiac symptoms, transthoracic echocardiographic examination revealed a large (33 mm × 42 mm) echo lucent cystic mass plugged into the interventricular septum. A positive serology test confirmed the diagnosis of hydatidosis. Treatment with antiprotozoal medication was started, and after two weeks, the patient successfully underwent surgical hydatidectomy using the cardiopulmonary bypass. This case highlights the rare concomitant cardiac, cervical, and hepatic involvement with hydatid disease. Together, a good understanding of the atypical manifestations, potential associated risk factors, and epidemiology leads to the optimal and timely management of such patients to minimize worse outcomes.

Case Presentation

A 68-year-old woman presented to a health care center with a clinical manifestation of a slow-growing and painless lump on the right side of the cervical region over several weeks. She had no cardiac symptoms. History-taking revealed working in a sheep-farming area in her twenties. Past medical and surgical history included hypertension and hepatic hydatidectomy two years ago.

Clinical examination through ultrasound imaging revealed a 47 mm × 75 mm cervical cyst expanded to superior mediastinum with neither inflammatory response nor spasm of the cervical muscles. The cervical cyst consisted of a bilayer membrane with several membrane-attached scolices, indicating an active hydatid cyst (cystic echinococcosis type 1, CE1). The lesion was lateral to the common carotid artery and posterior to the internal jugular vein with no cervical lymphadenopathy. Besides, the abdominal ultrasound examination showed multiple active, recurrent hepatic cysts in both the right and left lobes (stage 1), encompassing all liver segments. There was no evidence of biliary dilatation as well.

In the transthoracic echocardiographic (TTE), a bulging and well-defined echo-lucent cystic mass in the interventricular septum measuring 33 × 42 mm was detected (figure 1). A slight compression effect was present on the right ventricle (RV) cavity. The LV size and LV outflow tract (LVOT) were normal, with a mild systolic dysfunction (eye-ball estimation of LV ejection fraction = 45-5%). The valvular functions were normal, with no pericardial effusion. Other echocardiographic findings were unremarkable. The hydatid serology was positive, in which the enzyme-linked immunosorbent assay (ELISA)-based qualitative assessment of E. granulosus IgG antibodies confirmed the echinococcosis. Finally, the patient underwent cardiac surgery using cardiopulmonary bypass (CPB) for cystectomy to minimize the risk of spillage of cyst contents. The CPB technique was established by cannula inserting into the ascending aorta, superior vena cava (SVC), and inferior vena cava (IVC) after the routine median sternotomy. Following the cold cardioplegia, the established hypothermia was recorded at 32 °C. The outlines of the isolated cardiac cyst seemed to be complete and clear. Conservative procedures were further performed to sterilize and evacuate the cyst contents. The RV cavity was entered, and the cyst was exposed carefully. Thereafter, ten milliliters of its contents were aspirated.
An equal amount of hypertonic saline (NS 20%) was injected into the cyst, and after several minutes, the exposed cyst was evacuated completely (figure 2). Following successful excision and secured hemostasis, the cyst specimen, containing 8 ml colorless turbid fluid, was sent to the histopathological examination, which further vouched for the diagnosis of a hydatid cyst (figure 3).

Discussion

Due to well-developed transportation, echinococcosis, as a zoonotic disease, has become a serious global health problem, affecting more than one million people by hydatid disease worldwide [4]. Compared with visceral hydatidosis commonly occurring in the liver, cardiac HC is presented by wide clinical manifestations, leading to an early diagnosis challenge [13]. According to the WHO-Informal Working Group on Echinococcosis (WHO-IWGE) ultrasound classification, hydatid cyst consists of three stages, including active (CE1, CE2, with a high risk of rupture), transitional (CE3), and inactive or calcified cysts (CE4, CE5, with a low risk of rupture) [1]. Although most cardiac hydatic cysts in the literature are reported in young patients, here, we reported an old lady (68 y/o) with a recurrent hepatic hydatidosis accompanied by a huge homogenous cystic mass in the cervical and intracardiac regions. In our case, the cystic lesions in the vicinity of portal venous confluence and the left portal vein may be considered the leading cause of extra-hepatic hydatidosis. It is worth noting that the multiplicity and dispersion of the lesions, typical imaging findings, a history of husbandry procedures, a history of hepatic cysts, geographic location, and positive result of serologic test strongly established the hydatid cyst diagnosis. A previous study represented a 70 y/o female with no history of being in a sheep-rising area with signs in favor of right heart failure and cardiac hydatidosis complicated hydatid cyst and pre-tamponade [14]. Shojaei et al, in Iran, also indicated a cardiac HC in a 70-year-old farmer with dyspnea. The isolated lesion was diagnosed by echocardiography and further confirmed with cardiac MRI. Despite successful surgical excision, he died due to a progressive arrhythmia [15]. Another report in Iran has also documented an echinococcal infection involving an intramyocardial multicystic lump in the posterolateral and basal inferoseptal segments of LV in a 57-year-old farmer man referred with chest pain, and diagnosed by echocardiography, CMR, and positive ELISA-based serologic test. In contrast to our finding, EKG examination showed pathological Q and negative T waves. Similarly, surgery was the treatment of choice, followed by albendazole as a complementary therapy [3].

In conclusion, a good understanding of the atypical manifestations, potential associated risk factors, and epidemiology lead to the optimal and timely management of patients with rare echinococcosis to minimize worse outcomes.
Figure 1: Transthoracic echocardiography shows (A, D) in a four-chamber view a large, well-defined intramyocardial cystic lesion (with no obvious septation) in the mid part of the septum with mild compression effect on RVOT without gradient. (B, C) The parasternal long axis and short axis view revealed a cystic lesion.

Figure 2: (A) Surgery revealed a complete and clear cardiac cyst (B, C, D). After sterilizing and evacuating the cyst contents, the cardiac cyst was completely resected upon aspiration.
Figure 3: (A, B) Protoscolices of Echinococcus granulosus in cytology, ×400 magnification, Papanicolaou Stain.

References


