Abstract

Alveolar echinococcosis (AE), observed in the Northern Hemisphere, is caused by the larval stage of the fox tapeworm Echinococcus multilocularis. In endemic areas, annual incidence of AE ranges from 0.03 to 1.2/100,000 inhabitants. The liver is the primary focus of the disease but extrahepatic is also possible such as lung, spleen, pancreas, retroperitoneum, brain, bone and soft tissue. While pulmonary involvements occur in 7 to 20%, to our best knowledge; simultaneous intramyocardial and lung involvement is a very rare clinical entity reported in the literature.

Keywords

Alveolar Echinococcosis; Intramyocardial Involvement; Cardiopulmonary Surgery.

Özet

Alveolar kist hidatik (AKH) özellikle kuzey yarım kűrede etken Echinococcus Multilocularisın neden olduğu bir hastalıktır. Endemik bölgelerde insidansı 0.03-1.2/100 000 oranındadır. Karaciğer primer odak olarak bilinmesine rağmen akciğer, dalak, pankreas, retroperitoneal alan, beyin kemik ve yumuşak dokular da sık yerleştiği alanlardır. Pulmoner hidatik kist oransı % 7-20 olarak bilinmesine karşın eş zamanlı intramiyokardiyal ve akciğer tutulumu literatürde nadir görülen bir durumdur.

Anahtar Kelimeler

Alveolar Kist Hidatik; Intramiyokardiyal Tutulum; Kardiyopulmoner Cerrahi
**Introduction**

Alveolar echinococcosis (AE), observed in the Northern Hemisphere, is caused by the larval stage of the fox tapeworm Echinococcus multilocularis [1]. Human is infected after ingesting eggs, the metacestode cells of E. multilocularis which proliferates in the liver [2]. It is a potentially fatal, chronically progressive parasitic infection characterized by a long asymptomatic period and development of an invasive tumor-like lesion throughout this period [3, 4]. So early diagnosis of AE is very difficult because of long latent or asymptomatic period which may be as long as 15 years [2-4]. In endemic areas, annual incidence of AE ranges from 0.03 to 1.2/100,000 inhabitants [2].

The liver is the primary focus of the disease but extrahepatic involvement such as lung, spleen, pancreas, retroperitoneum, brain, bone and soft tissue. While pulmonary involvements occur in 7 to 20%, to our best knowledge; simultaneous intramyocardial and lung involvement sparing liver is a very rare clinical entity reported in the literature [5, 6].

Here, we present a very rare case of alveolar echinococcosis with simultaneous intramyocardial and lung involvement sparing liver which was completely removed at the same session via cardiopulmonary surgery.

**Case Report**

A 47-year-old woman was admitted to our hospital with a history of hydoptysis six months before admission. She didn't have any complaint otherwise. Physical examination did not reveal any abnormal findings: her lungs were normal on auscultation, no cardiac murmur or gallop rhythm was noted, and biochemical laboratory test results were within normal limits. Myocardial-specific enzyme values were within the normal range.

The patient's chest X-ray was normal, except for increase in convexity on left border of the heart; and the lesion paraatracheally located with sharp border, homogenous in content at upper zone of the left lung; and second one; sharp-bordered, paracardiac lesion with air-fluid content located at mid-zone of the left lung (Figure 1A). The Electrocardiography (ECG) showed normal sinus rhythm with T-wave inversion in leads I, aVL. (Figure 2A) transthoracic echocardiography showed a multivesicular cystic mass on apical part of the left posterolateral ventricular wall and protruded inward the ventricular cavity (Figure 2B,2C). The patient's abdominal ultrasonography and cranial computed tomography (CT) were normal. The patient's chest multislice CT and Magnetic Resonance Imaging (MRI) showed multivesicular cystic masses in the left ventricle lateral wall, at apical segment of the upper lobe of the left lung and at superior segment of the lower lobe of the left lung (Figure 1B-1D).

Serologically, specific Echinococcus granulosus and multilocularis antibodies were investigated by commercial ELISA (NovLisa, Echinococcus IgG Novatec, Germany For E. Granulosus; Bordier Affinity Products SA, Switzerland for E.multilocularis) and haemagglutination-inhibition test (HAI) (Hydatidose, Fumuoze laboratories, France) kits. Echinococcus multilocularis antibodies were positive with ELISA. Hydatid cyst antibodies was found to be positive 1/320 titer in the HAI.

The cardiac multivesicular cyst in the left ventricle wall was removed with cystotomy-capitonage procedure and then the cavity was washed with a hypertonic saline solution under cardiopulmonary bypass by conventional technique following median sternotomy. Gross appearance of the heart confirmed intramyocardial involvement (Figure 3A-D). Then, left pleural membrane was incised by using same median sternotomy and left lung was explorated for location of the pulmonary cysts, firstly the cyst located paravertebrally at left upperlobe apical segment, and then secondly the one located at left inferior lobe superior segment were removed with cystotomy-capitonage procedure and then the cavity was washed with a hypertonic saline solution by thoracic surgeon. The patient didn't experience any
complication early after operation and albendazole treatment was started immediately then after the surgery. The patient was discharged at sixth day postoperatively without any morbidity. And also she was under follow-up without any problem.

Discussion
Echinococcus multilocularis (E. multilocularis) is considered to be the most potentially lethal parasitic zoonosis in the nontropical areas in the Northern Hemisphere [6, 7]. Its invasive, tumor-like behaviour together with its long asymptomatic period to get clinical significance makes the alveolar echinococcosis potentially fatal disease. So early diagnosis is not possible in most of the cases. Cardiac echinococcosis is rare, representing only 0.5–2% of all cases [8]. Here, we reported a case of alveolar echinococcosis involving myocardium and lung sparing liver.

Chest pain, palpitations and dyspnea are the primary symptoms associated with cardiac echinococcosis [8, 9]. In our case those complaints never existed, so it supports also asymptomatic nature of the disease. That is so, whenever pulmonary cysts are in concern, exclusion of cardiac involvement should be always considered as we did in our case. But some clues can exist in ECG. In young patients from endemic areas, especially when the ECG shows a T-wave change, a diagnosis of cardiac echinococcosis should be suspected [8, 9]. In present case, T wave was negative at leads I, aVL. Since the patient didn’t have any risk factor for coronary atherosclerosis, further investigation was not performed.

Pulmonary AE is mainly caused by hematogenous dissemination from hepatic AE lesions. Physical signs and symptoms in pulmonary AE are hydoptysis, chest pain, cough with expectoration and exertional dyspnea. Hydoptysis is very typical of lung involvement. If present, prompt search for hydatid disease and to prevent recurrence of cysts after operation on lungs and heart [11]. The patient also was given albendazole immediately after the surgery.

In our case, there was presence of simultaneous intramyocardial and pulmonary multivesicular-type cyst hydatid. Through same median sternotomy and with single incision, the patient was operated for both cardiac and pulmonary cyst at same session to get complete cure.

In this case, we reported a very rare case of alveolar echinococcosis with simultaneous intramyocardial and lung involvement sparing liver and we aimed to share our clinical experience in the management of the case.

Competing interests
The authors declare that they have no competing interests.

References

How to cite this article: