A New Approach for Treatment of Bone Hydatid Cyst Disease: A Case Report of Primer Femur Hydatid Cyst

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INTRODUCTION

Hydatidosis is a tapeworm infection caused by the Echinococcus species (1). Involvement of the long tubular bones is rare in hydatid bone disease. Skeletal hydatidoses occurs in 0.5–2% cases (2). Bone hydatidosis with long bone involvement of the femur (16%) seen less commonly (3, 4). The cysts occasionally lie dormant in the body for up to 20 years without producing clinical signs or symptoms (3).

We report a case of hydatidosis of the femur with pathologic fracture in a 23-year-old man.

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CASE REPORT
A 23-year-old man presented to Emergency Department with a low energy femur fracture. He had fractured his femur while descending the stairs. Routine blood investigations was within normal limits except CRP was elevated [21.47 mg/dL] (normal was 0–5 mg/dL).

He has a history of operation from his left femur because of a fracture and a sliding hip plate was applied 10 years ago.

Radiographs of left femur revealed a multiloculated osteolytic lesions and subtrochanteric communucated fracture.with a sliding plate (Fig. 1).

![Fig: 1. Multiloculated osteolytic lesions and subtrochanteric communucated fracture.](image)

Computerized tomography (CT) show a multiloculated cystic lesion with cortical expansion and thinning of the cortices, extending from collum of femur to subtrochanteric region (Fig. 2). A low grade infection or malignancy was made for intial dignosis.
The patient was operated to for hardware removal and obtaining bone biopsy from the fracture side. In surgical gross anatomy cystic vesicules were seen in the fracture side. Histopathological examination revealed laminated hyaline membrane compatible with the diagnosis of hydatid cyst (Figs. 3, 4).

Fig: 2. Axial and coronal sections of tomography of patients.

Fig: 3. Histologic examination of the cyst wall shows an outer chitinous layer and an inner germinal layer which contains daughter cysts and brood capsules with scolices (Hematoxylin&Eosin x 40).
Blood test confirmed the hydatid cyst disease. An indirect hemiaglutinin test for echinococcal antibody was positive (1/1280) after the histopathological diagnosis was made.

After histopathological diagnosis was made magnetic resonance imaging was made and proximal 2/3 of entire femur was filled with hydatid cysts (Fig. 5).
Contamination of cysts are also seen between adductor muscles and vastus intermedius muscle. Because cyst hydatidosis mainly seen in liver and lungs, abdominal, thoracic and cranial CT are obtained and no abnormalities is seen.

Preoperative planning was made and tumour resection prothesis implantation desicion made. Because hypertonic solution were routine used in abdominal hydatidosis we also decide to use hypertonic saline solution. Patient operated and effected segments of femur (25 cm in length) removed and muscle planes are irrigated with hypertonic saline solution and a tumour resection prothesis (Mutars Total Femur Replacement Systems, Implantcast, Germany) is applied. Because of contamination of muscle mass a continuous negative pressure and irradiation wound closure system (VAC Instill® Therapy Unit, US) is used with irrigation solution of hypertonic saline for two weeks. After two weeks wound is closed in usual manner. A rehabilitation programme started and patient mobilized with crutches after two week.

The patients were administered postoperative chemotherapy with albendazole (10 mg/kg/day) for six months with close observation of the liver enzymes.

At the 12-month follow-up, patient is mobilized with only a cane and we detected no findings associated with local or systemic hydatid cyst. Radiological images taken after first year showed no recurrence and serological tests remained negative (Fig. 6).
Echinococcosis, which is known as unilocular hydatid disease or human hydatid disease is caused by the larvae of *E. Granulosus*. Bone hydatidosis commonly involves the spine and pelvis. Hydatid involvement of long tubular bones, especially the femur diaphysis, is rare (5, 6). The disease initially involves the epimetaphyseal region, often extending to the diaphyseal region at presentation (1). Progressive changes and an expanding cystic appearance may mimic tumour formation (7). In our case location was left femur extending from collum femoris to distal diaphysial area. A pathologic fracture had occurred before diagnosis in our case. Cystic appearance may be misdiagnosed as tumoural lesions so we perform a biopsy while hardware removal. It is difficult to diagnose bone hydatid cyst disease radiologically because the findings are non-specific (8). Plain radiographic findings in hydatid bone disease include intramedullary unilocular, bilocular and often multiloculated cystic expansile lesions with surrounding sclerosis in a honeycomb pattern associated with cortical
thinning (9). A definite diagnosis can only be reached by histopathological evaluation of resected tissues. Tuberculosis simple bone cyst, fibrous dysplasia, malign fibrous hystiosytom, sarcoma and metastasis should be consider in differantial diagnosis (10).

Surgical treatment is usually based on the same principles as described for a locally malignant lesion (1). The best treatment for bone infection originating from echinococcosis is removal of the involved bone or amputation (7). However, curettage and lavage with hypertonic salt, 1% formalin, or 0.5% silver nitrate solutions have been attempted (7).

Although wide excision of the infected bone provides the best chance of cure, it usually leaves an extensive bone defect that can produce a challenge during reconstruction (11). In our case we resected 2/3 of the femur and reconstruction of that was impossible.

Although amputation was one of the treatment alternatives, both using a tumour resection prosthesis with a continues irrigation with hypertonic saline solution gives us a change to save the limb. Recurrence of the disease can complicate prosthetic reconstruction after cyst excision (12). The current literature includes reports of patients whose hip was treated with total hip replacement. Two of these patients died due to complications; both died four years after surgery after chronic sepsis (2, 13). Neelapala et al, perform a revision hip replacement for a cyst hydatic but in his case extension of disease was through to ilium. Our case is the first case in litreture that treatment with a tumour resection prothesis with a continous irrigation with hypertonic solution.

As a conclusion hydatid disease involving the long tubular bones should initially be treated as a low-grade malignant neoplasm. In extensive lesions of bone and soft-tissue a tumour resection prosthesis with a continuos hypertonic saline irrigation can be an alternative treatment for amputation of the effected limb.
REFERENCES


