An Unusual Presentation of Hydatid Disease in a Child: A Case Report

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Abstract
The subcutaneous localization of the hydatid cyst in the abdominal wall is rarely encountered particularly in the pediatric population and is sometimes difficult to diagnose preoperatively. Here, we report the case of a 6-year-old boy who presented with two isolated episodes of low abundant hemoptysis and in whom a mass on the right lumbar region already considered as a lipoma was studied. Laboratory and radiological examinations were requested and the parietal cyst was surgically managed. The macroscopic and the pathological examination confirmed the diagnosis of the hydatid disease and helped in identifying the nature of the thoracic lesion that disappeared spontaneously after two episodes of hydatid vomiting. Hydatid cyst should be considered as a diagnosis for any masses of the abdominal wall. Moreover, biopsy and partial resection of the mass must be avoided.

Keywords
► hydatid cyst
► subcutaneous
► abdominal wall
► lung
► child
► case report

Echinococcosis or hydatid disease is defined as a parasitic zoonosis in which the Echinococcus cestode worms are the causative agents.1 There are two forms of the disease for which cystic echinococcosis (CE) and alveolar echinococcosis (AE) are pathogenic for humans.1 CE is caused by Echinococcus granulosus and is the most common presentation.1 The two most involved organs are the liver and the lung.2 The uncommon localizations such as soft tissues including abdominal walls are rarely seen even in endemic countries especially in children.2 Only five cases with similar localization in children were reported.3–7

We report a rare case of a child with subcutaneous hydatid cyst located in the lumbar region of the abdominal wall that its diagnosis was challenging in our clinical setting. The patient was successfully managed in our tertiary academic care center.

Case Report
A 6-year-old boy from a rural region was referred to our department of pediatric surgery by the family physician because of two isolated episodes of low abundant hemoptysis with an interval of 6 months. The child was afebrile and in a good general condition. There was no history of surgery, abdominal trauma, vomiting, abdominal or thoracic pain, hematologic disorders, or infectious diseases. Moreover, the patient had no family history of similar cases. Pulmonary vesicular sounds were decreased during auscultation of the right apical thoracic region. On the abdominal examination, a mass on the right lumbar region of 3×1.5 cm (cm) in diameter, firm in consistency, painless, and not fixed to the deeper tissue or to the overlying skin was found that was normal (►Fig. 1). It has evolved for 2 years and considered as lipoma at previous medical consultations. On hematological analysis, hemoglobin
was 12 g/dL, total leukocyte count was 9120/µL, eosinophil count was within the normal range, and C-reactive protein was 0.55 mg/L. All other biochemical serum parameters were without abnormalities. The hydatid serology was negative.

Chest X-ray showed a right middle pulmonary opacity (Fig. 2). Importantly, computed tomography (CT) scan of the chest revealed a fluid density lesion of the right hemithorax, heterogeneous, with irregular contours, not enhanced after injection of the contrast product with a periphery that is enhanced, measuring 39’40’28” mm in size (Fig. 3A and B). The abdominal CT scan supplemented by an ultrasound imaging showed a well-defined and anechoic cystic mass of the subcutaneous soft tissues of the right posterolateral abdominal wall that was enhanced in periphery after injection of the contrast product and measuring 32’23 mm in size (Fig. 4). The other abdominal organs were without any other abnormalities, particularly the liver, the spleen, the kidneys, and the pancreas. An anesthetic consultation was requested. The decision was to perform a surgical excision of the abdominal parietal mass under locoregional anesthesia that may help in diagnosis orientation. Surgical exploration showed a cystic mass of 4 cm in size containing a liquid and characterized by a “clear stone liquid” appearance and a proligerous membrane. The operating field was washed with a scolicidal agent using hypertonic saline solution (10% NaCl) and a complete excision of the cyst was performed (Fig. 5A and B). The macroscopic
The appearance of the mass was in favor of a hydatid cyst and the pathological examination of the specimens confirmed the diagnosis of the hydatid disease later, which helped in suggesting the nature of the pulmonary lesion. The child was discharged the same day of his admission in a satisfactory condition after an uneventful postoperative time. A medical treatment for 3 months was started immediately after surgery (albendazole 10 mg/kg/day). A thoracoscopy after 1 month was scheduled to manage the pulmonary localization, but two episodes of hydatid vomiting occurred before performing it leading to spontaneous disappearance of the pulmonary cyst (Fig. 6). No recurrence of the hydatid disease was detected after 3 months of follow-up and no negative incidents in terms of adherence and tolerability were observed. Taken together, the evolution of the disease in our patient, our management, and follow-up are represented in a timeline (Fig. 7).

Discussion

Hydatid disease remains endemic in Africa as well as in other regions worldwide particularly in South America, Mediterranean countries, Central Asia, and China. Morocco is one of the endemic countries of the hydatid disease with an overall prevalence of 1.9% in rural areas.

The liver is the most affected organ since it is the first and the largest filter of parasitic embryos coming from the intestine through the portal system. The lungs and rarely the other organs are also exposed by the systemic circulation after possible escape from this filter. Multiple organs in hydatid cyst are involved in 10.3% of cases and the association liver-lung is described to be the most occurred (7.7%). It was reported that the incidence of hydatid cysts with uncommon localization represents globally 7% and the cases involving soft tissue reach only 1% in pediatric population. This low prevalence can be explained by muscular contracility and the presence of lactic acid making the growing of the cyst difficult. To the best of our knowledge, only five cases of hydatid cysts involving the abdominal wall in
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children were reported. Those cases interested the right lower abdominal wall, the suprapubic region, the right inguinal, and the lumbar regions.

The hydatid cyst is generally asymptomatic or minimally symptomatic. For this reason, it can reach large dimensions. Clinical manifestations are nonspecific and depend on the location, the size of the cyst, and the pressure effect of the slow-growing cyst in the affected organ. In soft tissues, it is usually described by the clinicians as painless and noninflammatory swelling, which is consistent with the clinical course of our patient.

Serological testing can confirm the diagnosis, but its accuracy is not high. The localization of the hydatid cyst can clearly influence the intensity of the serological response of the body. Notably, serology of hydatid disease can be positive in most cases but only in 27% of patients with musculoskeletal location. In addition, a significant association between a serological negative test and the rupture of the cyst was previously reported. The hydatid serology of our patient was negative that is in line with previous literature findings.

Chest X-ray, ultrasound, CT, and magnetic resonance imaging may be necessary depending on the organ affected and the stage of the disease to confirm the diagnosis. However, an atypical localization of the hydatid cyst with inconclusive imaging findings can be challenging. The hydatid cyst of the soft tissue could be seen radiologically under various aspects including unilocular cyst, cyst with detached membranes, with daughter cysts or with calcified wall. In our case, the mass was seen by ultrasound and CT as unilocular cyst without detached membranes, daughter cysts, or calcified wall making the diagnosis not evident.

Lipoma, which was the diagnosis considered before the presentation of the patient to our department, the sebaceous cyst, and abscess might mimic a hydatid cyst of soft tissues. For this reason, the hydatid disease should always be kept in mind especially in endemic areas to avoid biopsy or partial excision of the mass. Otherwise, the risk of anaphylaxis or recurrence of the disease after spillage should be considered.

The available therapeutic options to clinicians in term of soft tissue hydatid disease include surgery, chemotherapy, and puncture/aspiration/injection/reaspiration (PAIR). Surgical cystectomy is the basis of the treatment. The goals are to remove completely the cyst and to avoid further contamination caused by spillage of cyst content. In our patient, an accidental opening of the cyst during the dissection was occurred and the contamination was avoided by washing the operating field with sclocial compounds based on hypertonic saline solution. Until to date, the medical treatment based on benzimidazole compounds (albendazole and mebendazole) is still useful to prevent the recurrence and to reduce the size of the hydatid cysts. Importantly, a previous report by Srivastava et al showed in a case of a 14-year-old boy with a hydatid cyst of the right lower abdominal wall, of 11.2 x 3.5 cm in size, with multiple daughter cysts that medical treatment using albendazole as monotherapy may also have a place in the therapeutic strategy of the hydatid disease in addition to surgery. In fact, the cyst disappeared after 14 weeks following the albendazole treat-

ment alone. Furthermore, PAIR can be used for inoperable patients or for those who are reluctant to surgery. However, it is necessary to be accompanied by a medical treatment. In our case, the diagnosis of hydatid disease was considered intraoperatively and the medical treatment was started immediately after surgery. During his last follow-up visit, our patient achieved a complete remission and his family was satisfied with our management strategy.

The diagnosis of hydatid cyst with unusual localization remains challenging even in endemic areas of the disease, but it should be kept in mind to avoid any biopsy or partial resection. A total surgical resection of the cyst is the treatment of choice, which should be combined to benzimidazole drugs to prevent possible recurrence. Other localizations particularly in the liver and lungs should be systematically searched.

Statement of Ethics
Written informed consent was obtained from the child’s parents to publish this case report and all associated images. A copy of this written consent is provided if requested by the journal/editor.

Author Contributions
K.E collected patient’s data, conducted the literature review, and wrote the manuscript. A.A, I.K and H.B reviewed and supervised the case report writing. All authors approved the final version of this paper.

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Conflict of Interest Statement
The authors have no conflicts of interest to declare.

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