CASE REPORT

Pancreatic hydatid cyst mimicking a choledochal cyst

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Abstract
Hydatid disease is endemic to South Africa. Radiological findings may help establish the diagnosis and aid preoperative preparation. Unusual locations and atypical imaging appearances may complicate the differential diagnosis. A case of isolated hydatid disease mimicking choledochal cyst is presented. Primary hydatid disease of the pancreas in the paediatric population is rare and this is only the sixth case reported in the English literature. This article highlights the importance of always considering hydatid disease in our patient population whenever a cystic lesion of the pancreas is identified.

Introduction
Hydatid disease is a parasitic infection caused by the larval stage of the tapeworm Echinococcus. Human infection is endemic in sheep-raising areas of the world, including South Africa. Pancreatic involvement in adults occurs with a reported prevalence of 0.25%. Primary pancreatic hydatid disease in childhood with no other associated visceral involvement is extremely rare, and only 5 cases have been reported to date. Nonetheless, hydatid disease should always be included in the differential diagnosis when a cystic lesion is identified in a patient from an endemic area.

Case report
An 11-year-old girl was admitted with symptoms of epigastric pain and vomiting. Examination revealed a palpable epigastric mass and jaundice. The latter was confirmed biochemically. The serum amylase was normal. Abdominal ultrasound (Fig. 1a and b) and computed tomography (CT) scan (Fig. 2a and b) of the abdomen were obtained.

The differential diagnosis considered included choledochal cyst, congenital cyst, pseudocyst, cystadenoma and hydatid disease. In view of the symptoms, the patient was operated on and the cyst removed. Histopathological examination revealed flimsy translucent membranous material confirmed on microscopy to be consistent with echinococcal disease.

Discussion
Pancreatic cysts are uncommon in children. Cysts may be congenital (i.e. developmental) or acquired. The latter includes pseudocysts, retention cysts, duplication cysts, neoplastic cysts and parasitic cysts. Choledochal cysts, although not arising from the pancreas, are intimately related and may sometimes be difficult to distinguish from pancreatic cysts on imaging studies.

Hydatid disease in humans begins with ingestion of eggs, followed by penetration of the gut wall by the embryo and entrance
It is postulated that the cyst wall may be a thin well-laminated membrane and the thin inner germinal membrane. A type II cyst has daughter cysts and/or matrix. When the hydatid fluid is replaced by formed elements the lesion dies and eventually becomes calcified and biologically inert, forming a type III lesion. Complications of hydatid disease including rupture and secondary infection are classified as type IV cysts. Three subtypes of rupture are described, viz. contained rupture, communicating rupture and direct rupture with peritoneal spill. On ultrasound, the appearance of visceral hydatid disease varies according to the stage. The cyst wall may be a thin well-defined echogenic rim but in certain cases double echogenic lines separated by a central hypoechoic layer have been described. Internal architecture is also variable. In this case a unicellular anechoic lesion was seen but internal septa, floating membranes, daughter cysts and echogenic hydatid sand have also been described. The latter creates a snowstorm appearance when the patient is turned over and scanned immediately. Multilocular cysts may manifest with a cyst within a cyst appearance or a honeycomb pattern with multiple septa, each septum representing the wall of the daughter cyst. Curvilinear cyst wall and internal calcification is also seen as a hyper-echoic contour with acoustic shadowing.

CT may display the same findings as ultrasound. The cyst fluid is usually of water attenuation unless complicated by super-added infection. The cyst wall is usually well defined except in super-added infection, with variable nonspecific contrast enhancement. The classic ‘wheel spoke’ appearance described on ultrasound is also seen on CT scanning (Fig. 3).

Review of the 5 previously described paediatric pancreatic hydatid cysts, confirmed unicellular cysts with no specific distinguishing features in 3 instances. The other 2 cases had more characteristic type II features. In conclusion, the appearance of a cystic mass with an undulating membrane, as with contained rupture, or of multiple degenerating daughter cysts within the mother cyst may alert the clinician to the possibility of pancreatic hydatid disease.

On the other hand, radiologic examinations alone may not be sufficient to diagnose primary pancreatic hydatid disease and serological tests may be necessary. In our case, both US and CT were useful in diagnosing the cystic mass in the head of the pancreas, but were unhelpful in lesion characterisation. The radiologist should always strive to make the diagnosis of hydatid disease as direct rupture has great clinical significance, including anaphylaxis, hydatid dissemination and secondary bacterial infection.

References