Cystic lymphangioma of the pancreas

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Summary
Cystic lymphangioma of the pancreas is a rare condition. A 14-year-old girl presented with a cystic abdominal mass and abdominal pain. She was initially treated by biopsy and cyst drainage, and subsequently with partial excision of the cystic mass. The mass, which was a cystic lymphangioma involving the pancreas, recurred after each operation. Persisting symptoms led to a pylorus-preserving pancreatectomy and hepaticejunostomy. Total excision is the only effective method of treating this benign tumour. The patient remains symptom free 2½ years after pancreatectomy.

Keywords: cystic lymphangioma, pancreas, abdominal mass

Intra-abdominal cystic lymphangiomas involving the mesentery, omentum, and retroperitoneum are rare benign tumours. Reported incidence rates vary from 1 in 25 000 to 1 in 250 000 hospital admissions. Cystic lymphangioma involving the pancreas is rarer still, only eight cases having been reported in the last 10 years. The majority of cystic lymphangiomas are discovered in childhood as asymptomatic abdominal masses.

Case report
A 14-year-old girl presented to a paediatric surgeon with an abdominal mass. She had complained of vague abdominal pains for the last year and on examination was found to have a swollen abdomen with a central abdominal mass. An ultrasound scan showed a large mixed solid and cystic mass, that extended from the left lobe of the liver to the splenorenal angle, lying anterior to the inferior vena cava and aorta. The pancreas could not be identified separately from the mass. The differential diagnosis suggested was a tumour of hepatic or ovarian origin.

Computed tomography (CT) showed that the head of the pancreas was involved by the mass and a diagnosis of cystadenoma of the pancreas was considered (figure 1). An exploratory mini-laparotomy in January 1992 revealed an haemangiomatous lesion of the pancreas and a blood-filled cyst. This was biopsied and drained. Histological examination showed features consistent with a benign lymphangioma. In May 1992, due to persistent enlargement of her abdominal mass, laparotomy and partial excision of the tumour was undertaken.

During follow-up over several months her symptoms of abdominal discomfort and postprandial fullness returned. Repeated ultrasonography confirmed continued enlargement of the mass and its origin from the pancreas. She was referred to us for further management.

At a third laparotomy in December 1992 extensive cysts were found in the lesser sac and portahepatis encircling the duodenum and replacing the pancreatic head. Trial dissection to separate the mass from the pancreas failed. A pylorus-preserving pancreatectomy with hepaticejunostomy was performed. Macroscopically, the specimen consisted of a multicystic, multilocular mass, 14 cm in diameter, diffusely expanding the body of the pancreas. Microscopy showed variably sized, massively distended, lymphatic channels with apparent origin within the pancreas (figure 2). The diagnosis was of diffuse cystic lymphangioma of the pancreas.

Two and a half years later she remains well, with normal weight gain and development, with no evidence of further symptomatic cysts.

Discussion
Wegner classified lymphangiomas by histological appearance in 1877. He separated these tumours into simple, cystic and cavernous lymphangiomas. Cystic lymphangiomas are classified as benign tumours but their exact origin is unknown. Many theories have been formed to account for their development, including increased production or retention of lymphatic fluid, and infection or inflammation of the lymphatic channels. A developmental abnormality has also been blamed and this has led to cystic lymphangioma being classified as a hamartoma rather than a true neoplasm. Malignant transformations to lymphosarcoma or adenocarcinoma have been documented but are exceedingly rare.

Intra-abdominal cystic lymphangiomas may be multiple or single, unilocular or multilocular. It is believed that these different manifestations depend on whether there is sequestration or obstruction of the lymphatic vessels. If obstruction predominates then large unilocular solitary omental or mesenteric cysts tend to develop. If budding or sequestration of the lymphatic channels occurs then a typical cystic lymphangioma will develop that is multilocular and may be present in more than one site. At operation it can sometimes be difficult to differentiate mesenteric cysts from cystic lymphangioma. The correct diagnosis can be obtained from histological examination. It is important to obtain the correct histological
Cystic lymphangioma of the pancreas

Figure 1 (A) CT of the abdomen without intravenous contrast injection showing a large mixed solid and predominantly cystic mass (upper arrow) involving the pancreas (lower arrow). (B) The mass extends to the right side of the pelvis and distends the anterior abdominal wall.

Figure 2 (A) Low-power microscopy to show large ectatic lymphatic channels separating lobules of pancreas, H & E x 40. (B) Pancreatic acinar tissue (bottom right) and dilated vascular channels lined by endothelium, with lymphoid aggregates in the stroma of the walls (H & E x 100).

diagnosis because these two entities behave in completely different manners: cystic lymphangiomas are more aggressive and will invade other organs and hence the chances of complications or recurrence after excision are greater than for a mesenteric cyst.

Cystic lymphangioma may occur at numerous sites throughout the body (box 1) but less than 1% are intra-abdominal. Retroperitoneal cystic lymphangiomas are usually a condition of childhood, 90% presenting before the age of two. The contents of the cysts can vary from serous to purulent, bloody to chylous. No difference in geographic distribution or sex predilection has been identified.

The clinical presentation of these childhood tumours is varied. Small cystic lymphangiomas will remain asymptomatic and undetected. Larger growths may be detected incidentally on abdominal examination for an unrelated complaint. As growth of the mass occurs the patient may complain of abdominal distension and pain. Acute symptoms suggest pressure on neighbouring structures or peritonitis as a result of rupture, torsion, haemorrhage, or infection of the cyst. The commonest presentation is that of an asymptomatic abdominal mass.

There is a wide differential diagnosis for children presenting with an abdominal mass. In the acute setting the commonest diagnosis made is that of acute appendicitis. Even with the aid of pre-operative investigation, the diagnosis of cystic lymphangioma is seldom made. Ultrasonography is the initial investigation of choice but, despite high sensitivity at detecting cystic lesions (94%), the correct diagnosis will be suspected in less than 25% of patients. After CT and fine needle aspiration cytology the diagnosis is more likely to be considered. Magnetic resonance imaging (MRI) is equal to CT in demonstrating these cystic growths and avoids exposure to ionising radiation.

Sites of distribution of cystic lymphangiomas

<table>
<thead>
<tr>
<th>Location</th>
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<tr>
<td>75% neck</td>
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<tr>
<td>20% axilla</td>
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<tr>
<td>4% mediastinum, lungs, chest wall, arms, back, parotid, spleen, liver, pelvis, groin and skin</td>
</tr>
<tr>
<td>1% retroperitoneum and omentum</td>
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Box 1

Differential diagnosis of cystic abdominal masses in children

- ovarian cystadenoma or teratoma
- pancreatic cystadenoma
- renal: simple and multilocular cysts, polycystic kidney disease, cystic nephroma, hydronephrosis
- liver: choledochal cyst, cystic hepatoblastoma, hydatid cyst
- splenic cyst
- periappendiceal abscess
- intramuscular
- enteric cyst or duplication
- mesenteric lipoma
- retroperitoneal neoplasm
- ascites

Box 2
The treatment of choice is total excision. If there has been invasion of other tissues then complete excision including the affected area of the involved organ is necessary. Partial resection is associated with a high recurrence rate, 50% in one series after a mean follow-up period of two years. After complete excision recurrences can occur but the rate is much lower at 7%. Aspiration, cystenterostomy and peritoneal cavity marsupialization are usually followed by early recurrence.

Our case demonstrates the difficulty of diagnosing these lesions pre-operatively and the risk of recurrence following simple drainage or partial excision. Ultrasonography will confirm the cystic nature of the mass, however, the correct diagnosis will only be reached if intra-abdominal cystic lymphangioma is considered in the differential diagnosis of abdominal cysts. CT or MRI is necessary to delineate the extent of the tumour and to identify involvement of any other organs prior to planning surgery. Complete excision is essential to prevent an unnecessarily high recurrence rate.

Pancreatic cystic lymphangiomas probably arise from extension of a retroperitoneal cystic lymphangioma anteriorly into the pancreas. They represent the severe end of the spectrum of intra-abdominal cysts. Our patient was atypical in that she presented at age 14 with a symptomatic mass. The correct diagnosis was not made before initial laparotomy and biopsy. Drainage and subsequent partial excision of the cystic lymphangioma were soon followed by recurrence of the mass. She was treated successfully by partial pancreatectomy and total excision of the mass and she remains free of symptomatic recurrence 30 months after surgery.

Learning points

- cystic lymphangioma of the pancreas is a rare, benign, tumour
- they usually present in infancy, 90% occurring before the age of two
- an asymptomatic abdominal mass is the commonest presentation
- acute symptoms are uncommon and tend to be misdiagnosed as appendicitis
- ultrasound combined with CT or MRI aids diagnosis and planning of surgery
- complete excision is essential to prevent recurrence

Box 3

Cystic lymphangioma of the pancreas.

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