CASE REPORT

CHOLEDODHAL CYST AND CHRONIC PANCREATITIS — TREATED BY PROXIMAL PANCREATECTOMY

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A 32-year-old woman with a choledochal cyst (Todani type I) developed recurrent acute pancreatitis leading to calcific chronic pancreatitis. She had previously been treated with two cyst drainage procedures and subtotal cyst excision. This association between choledochal cyst and chronic pancreatitis has not been previously reported. Severe continuing symptoms led to pylorus-preserving proximal pancreateodudeneectomy, which was undertaken to prevent future carcinoma in the cyst remnant and progression of the chronic pancreatitis.

KEY WORDS: Choledochal cyst, chronic pancreatitis, pylorus preserving proximal pancreateoduodenectomy

INTRODUCTION

The incidence of choledochal cyst in Western countries is estimated at one in every 100,000–150,000 live births1. There is a female preponderence of nearly 4:12. Complications of choledochal cyst in adult life include gallstones, cholangitis, cholangiocarcinoma and pancreatitis3,4. In Yamaguchi’s collected series of 1433 patients from Japan, there were six cases of pancreatitis of unspecified type5. We have recently reported two patients with recurrent acute pancreatitis4, and this association is well recognised though uncommon2,6,7. Although chronic calcific pancreatitis can be the end stage of recurrent acute attacks8, it does not appear to have been reported as a complication of choledochal cyst. We now describe such an association in a young woman.

CASE REPORT

A 32-year-old woman was referred to this hospital after four previous operations for choledochal cyst. At the age of 3 months and again at 12 years, she had undergone a cyst drainage operation. Thereafter she was well until the age of 30
years, when repeated episodes of cholangitis led to subtotal excision of the cyst with Roux-en-Y hepaticojejunostomy. Soon afterwards attacks of acute pancreatitis commenced, each time with marked elevation of serum amylase. An endoscopic retrograde cholangiopancreatogram (ERCP) showed a 5mm calculus in the pancreatic duct in the head of pancreas, and further operation was undertaken. Despite a wide transduodenal sphincteroplasty, the orifice of the pancreatic duct could not be reached nor could the stone be extracted. By the time of referral to the Hammersmith Hospital, she had had at least four further attacks of documented acute pancreatitis plus numerous minor episodes. Alcohol intake had never exceeded 3-4 units per week nor had gallstones ever been found.

Normal investigations included liver function tests, serum amylase, glucose tolerance test and pancreolauryl ratio for exocrine pancreatic function. Computed tomography showed dilatation of the residual intrapancreatic bile duct and a focus of calcification in the pancreatic head. Repeat ERCP confirmed a sizeable remnant of choledochal cyst plus two calculi partly obstructing the pancreatic duct (Figure 1a,b).

Resection of the pancreatic head seemed the best approach for this complex problem. A 3 cm remnant of the choledochal cyst was dissected free from the Roux loop leading to the biliary anastomosis and from the hepatic artery and portal vein. Following pylorus-preserving proximal pancreaticoduodenectomy, a retrograde pancreaticogram from the neck showed dilatation of the distal pancreatic duct, but the body and tail of pancreas appeared much less severely diseased than the head. The previous hepaticojejunostomy was not disturbed, so reconstruction involved end-to-end pancreaticojejunostomy and end-to-side duodenojejunostomy (Figure 2a,b). Recovery was uneventful and postoperative glucose tolerance test was unchanged. Since the pancreolauryl ratio had fallen from 35% to 7% (normal >30%), pancreatic enzyme therapy was started postoperatively. Nine months later she is well and free of pain.

The resected specimen showed an anomalous pancreaticobiliary ductal junction (PBDJ). The common channel measured 20 mm in length. It received an inflamed, dilated main pancreatic duct containing the two stones and, via a pin-point orifice, the remaining portion of the choledochal cyst. The cyst showed no evidence of malignancy. There was clear-cut chronic pancreatitis in the head of pancreas with focal areas of active inflammation.

DISCUSSION

The association between cystic dilatation of the biliary tree and an anomalous PBDJ has been clearly established by Japanese workers. The abnormal common channel, which is >15 mm long instead of the normal <5 mm, probably allows reciprocal reflux of bile and pancreatic juices. Amylase can be shown in the bile and may explain the increased incidence of cholangiocarcinoma and gallbladder carcinoma. As a corollary, entry of bile into the pancreatic duct might lead to recurrent acute pancreatitis with the eventual development of chronic inflammation, fibrosis and calcification of the pancreas. Surprisingly, we can find no previous report of a patient with choledochal cyst and calcific chronic pancreatitis.

Excision of a choledochal cyst is nearly always preferable to simple drainage, which runs the risk of recurrent cholangitis and carcinoma. In our patient, the
Figure 1a  ERCP showing lobulated remnant choledochal cyst.

Figure 1b  Dilated pancreatic duct with calculus (arrow).
Figure 2a,b Reconstruction following pylorus preserving proximal pancreatoduodenectomy.
cyst excision performed 17 years after the second drainage procedure succeeded in preventing further cholangitis but seemed to precipitate the development of acute-on-chronic pancreatitis. Perhaps the intrapancreatic cyst remnant caused stasis of pancreatic juice and led to calculus formation in the duct. The extent of chronic pancreatitis is manifested by the fact that a limited pancreatectomy (40–50 per cent) has led to near-total exocrine insufficiency.

In Todani’s series of 73 patients with excision of choledochal cyst, seven patients needed reoperation for bleeding, cholangitis or anastomotic leakage. Two patients developed acute pancreatitis but neither required reoperation. Thus, pancreatitis represented a unique indication for reoperation after cystectomy in our patient. By removing the cyst remnant and the obstructing pancreatic calculi, conservative proximal pancreatectomy should prevent future carcinoma and progression of chronic pancreatitis.

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INVITED COMMENTARY

The association between attacks of acute pancreatitis and the presence of a choledochal cyst is now well recognized and is usually related to the presence of a 'common' pancreatico-biliary channel at the ampulla. Babbitt (1969) first described common channel between the lower end of the bile duct and the pancreatic duct in 19 cases of choledochal cyst and he suggested that pancreatic reflux might damage the common bile duct and cause the dilatation. These anomalous junctions have now been described in many reports and in a personal series of 39 cases no less than 29 had long common channels and high levels of amylase within the bile.

The pancreatitis in these patients should be differentiated from so-called fictitious pancreatitis. Free reflux of pancreatic juice can occur through any common pancreatico-biliary channel and amylase may be absorbed through the choledocho-cyst wall which usually has an incomplete epithelial lining. Hyper-amylasaemia may therefore be detected without the inflammation of the gland. We have also seen recurrent pancreatitis in 3 out of 4 cases treated with cyst enterostomy, rather than radical cyst excision, and all of these had a common pancreatico-biliary channel.

Diversion of bile by cyst excision and Roux loop reconstruction usually prevents any further episodes of pancreatitis. The present case report is therefore unusual. Firstly, because bile drainage had been diverted previously and secondly because the patient developed a chronic form of pancreatitis. There was however partial obstruction by calculi in the pancreatic duct and a retained portion of choledocho-cyst behind the duodenum which may have harboured infection. Both of these factors have been implicated in the aetiology of pancreatitis (Schmidt and Creuzfeldt, 1976).

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