Granular cell tumour of the bile duct in association with intrahepatic bile duct adenomas

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Granular cell tumour of the bile duct in association with intrahepatic bile duct adenomas. Can J Gastroenterol 1994;8(2):92-96. Granular cell tumour of the extrahepatic biliary tract is a rare benign lesion likely of neurogenic origin. Review of the previously reported cases indicates that almost all patients are female, and the majority is Black. Symptoms usually are those of biliary obstruction or cholecystitis. Surgical resection of the tumour is curative. Intrahepatic bile duct adenoma is another rare benign biliary neoplasm that does not manifest clinically but can be confused with metastatic carcinoma, cholangiocarcinoma or other focal liver lesions at laparotomy or autopsy. The authors report the case of a symptomatic Caucasian woman with biochemical evidence of liver disease who had a granular cell tumour of the bile duct as well as several intrahepatic bile duct adenomas.

Key Words: Bile duct tumour, Granular cell tumour, Intrahepatic bile duct adenoma

Granulome du canal cholédoque, en association avec un adénome intrahépatique : Rapport de cas et survol de la littérature

RÉSUMÉ : Le granulome du tractus biliaire extra-hépatique est une lésion bénigne susceptible d’avoir une origine neurogène. Le survol des cas recensés à ce jour indique que la plupart des patients sont de sexe féminin et en majorité de race noire. Les symptômes sont habituellement ceux d’une obstruction bilaire ou d’une cholecystite. L’ablation chirurgicale de la lésion est curative. Le granulome du tractus biliaire extra-hépatique est une autre néoplasie biliaire bénigne rare qui n’a pas de manifestations cliniques, mais qui peut être confondue avec un cancer métastatique ou avec un cholangiocarcinome ou une autre lésion hépatique localisée à la laparotomie ou à l’autopsie. Les auteurs rapportent le cas d’une femme de race blanche asymptomatique manifestant des signes biochimiques de maladie hépatique qui présentait un granulome au niveau du canal cholédoque et plusieurs adénomes des voies biliaires intrahépatiques.

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Received for publication July 26, 1993. Accepted August 16, 1993

EXTRAHEPATIC BILIARY OBSTRUCTION is most often caused by gallstones or malignant tumours. Benign neoplasms of the extrahepatic biliary tree are very uncommon, with the majority being adenomas and papillomas (1).

Granular cell tumour (GCT) involving the biliary tree is exceedingly rare; fewer than 50 cases were reported in the literature up to 1992 (2-7). GCTs usually cause biliary colic or jaundice and have a predilection for young Black females.

We report the case of a young Caucasian woman who, in the absence of biliary symptoms, was found to have a GCT of the bile duct as well as intrahepatic bile duct adenomas, another form of rare benign biliary neoplasm.

CASE PRESENTATION

A 34-year-old previously healthy Caucasian woman presented to the emergency department with a three-year history of tension headaches for which she took up to 25 tablets of acetaminophen (325 mg each) per day. She denied any other symptoms such as abdominal pain, jaundice, fever, pruritus or malaise. The physical examination was completely normal. She was prescribed chloral hydrate for her headaches with prompt relief.
A biochemical screen showed the following abnormalities (normal range in brackets): alkaline phosphatase 451 U/L (39 to 117); aspartate aminotransferase 73 U/L (7 to 32); alanine aminotransferase 143 U/L (0 to 31); and gamma-glutamyl transferase 702 U/L (7 to 32). Total bilirubin was 5 µmol/L (normal 4 to 18). An abdominal ultrasound revealed a normal gallbladder and prominent intrahepatic bile ducts. The common duct measured 0.9 cm in diameter. A solid echogenic lesion, 1.5 cm in width, was noted to be within the common bile duct (Figure 1).

Endoscopic retrograde cholangiopancreatography demonstrated pancreas divisum and suggested a biliary stricture, which was more clearly demonstrated on a percutaneous transhepatic cholangiogram. The stricture was located at the junction of the common hepatic duct and the cystic duct. The distal common bile duct was normal but there was mild dilation of the biliary radicals proximal to the stricture (Figure 2). At surgery there was a hard 1.5 cm nodule with surrounding fibrosis just above the junction of the cystic and common bile ducts. Several small, raised lesions, 3 to 4 mm in diameter, were noted on the liver surface. These grossly resembled metastatic deposits and were excised. The pancreas was normal. The tumour was resected, cholecystectomy was performed and the biliary tree was reconstructed with a choledochoduodenostomy. The patient's subsequent clinical recovery was uneventful.
PATHOLOGICAL FINDINGS
A poorly defined yellowish mass measuring 2x1 cm was present in the wall of the common bile duct close to the entry of the cystic duct. The proximal resection end of the bile duct was dilated. The gallbladder was normal. Two wedges of liver tissue, each measuring 1x1 cm, were mostly replaced by well-circumscribed nodules.

Microscopic examination revealed infiltration of the whole thickness of the common bile duct wall by strands and solid clumps of large eosinophilic finely granular cells with vesicular nuclei (Figure 3). The tumour cells stained positively for S-100 protein and polyclonal neuron-specific enolase using avidin-biotin immunoperoxidase technique. Electron microscopic examination demonstrated membrane-bound autophagic granules and distinct basal lamina (Figure 4).

Microscopic examination of the excised surface lesions of the liver revealed a network of ductal structures embedded in a fibrous stroma (Figure 5). The small and inapparent lumina were lined by low columnar epithelial cells lacking nuclear pleomorphism. There was no evidence of bile within the ductules. A mild lymphocytic infiltrate was present. The adjacent liver parenchyma was unremarkable.

DISCUSSION
GCT is a rare neoplasm found usually in the oral cavity, dermis and subcutaneous tissues of the extremities and chest wall, though it can occur in virtually any organ system (8). Gastrointestinal involvement occurs in less than 5% of GCTs, may be multicentric and usually is asymptomatic due to its submucosal location (9).

GCTs of the biliary tract, first observed by Coggins (10), have been previously recorded in 45 patients; Butterly et al (2) reviewed 39 patients and six additional cases have appeared in the literature (3-7). Mean age at presentation is 33.6 years (range 11 to 61) for all cases. Twenty-nine of the 45 patients (64%) were Black and only three were males (93% female incidence). A greater than expected frequency of GCTs - both biliary and extrabiliary - among Black patients has been reported previously (2,8), and the marked female predominance in cases of biliary involvement is striking.

Approximately half of the GCTs is located at or near the junction of the cystic duct, common bile duct and common hepatic ducts. More proximal hepatic duct involvement has been described in only two patients (2,11). In 14 cases, the tumour was confined exclusively to the cystic duct (2,3), and in two instances, isolated gallbladder lesions were seen (5,12). Only one patient had multifocal tumours in the common bile duct, cystic duct and gallbladder (13).

The clinical manifestations generally reflect the location of the tumour. Common bile duct lesions present with pain and/or jaundice in more than 90% of patients. Pruritus was the initial complaint in a few patients (6,7,14-17). When the cystic duct or gallbladder was affected, recurrent biliary colic or acute cholecystitis occurred (2). In one patient, a common bile duct GCT was found incidentally at autopsy (18).

This report describes the first living asymptomatic patient with such a le-
tion (discovered because of an abnormal liver enzyme profile and ultrasound). It is likely that further expansion of the tumour would have led to symptoms pointing towards the biliary tree.

As in the presented case, GCT of the biliary system is an unexpected finding intraoperatively after ultrasound and cholangiography have demonstrated an obstructing lesion or stricture. Cholangiocarcinoma, sclerosing cholangitis or stone disease is the usual preoperative diagnosis.

The tumours grossly are yellow-white in colour, usually less than 3 cm in diameter, nodular and not encapsulated. On light microscopy, GCTs consist of large polygonal cells arranged in small nests and cords, and are associated with a variable amount of fibrosis. The round nuclei are centrally located and the eosinophilic cytoplasm contains multiple coarse granules that stain positively with periodic acid-Schiff stain (6).

Abrikossoff (19) initially coined the term ‘granular cell myoblastoma’, postulating a histogenetic origin from striated muscle. However, current knowledge favours Schwann cell derivation. Supporting evidence includes the tumours’ close association with nerves and shared immunohistochemical reactivity with antibodies directed against several neural proteins, such as S-100 protein, neuron-specific enolase and myelin-basic protein. Electron microscopic demonstration of neural elements also lends support to the Schwann cell theory (6). Nevertheless, the noncommitted term ‘granular cell tumour’ remains widely accepted.

The treatment of biliary GCTs is surgical resection and reanastomosis. Buttery et al (2) observed that in almost all instances these tumours are located in an area where curative excision is readily accomplished and reconstructive procedures – including choledochoduodenostomy, choledochojejunostomy, hepaticoduodenostomy and hepatojejunostomy – can be carried out. Intrapancreatic common bile duct involvement may require more extensive surgery, i.e., pancreaticoduodenectomy or a simple bypass procedure depending on the clinical circumstances. So far there have been no recurrences after surgical resection of biliary GCTs in all reported cases. Malignant GCTs have been described previously (20), but there are no reports of such tumours involving the biliary tract. Concurrent GCTs at extrabiliary sites have been observed in six patients (4, 7, 11, 21, 22). In a recently reported patient with small bowel mesentry and tracheal involvement in addition to a GCT of the bile duct flow cytometric DNA analysis showed that all lesions were diploid with low proliferation, supporting multicentricity rather than metastasis (7).

Several intrahepatic bile duct adenomas were discovered in the presented patient intraoperatively. These tumours are exceedingly rare; in a series of 50,000 necropsies only four cases were identified (23). They are more prevalent in males than females, with two-thirds of the reported patients older than age 50. These lesions are virtually always incidental findings during laparotomy or autopsy (24). Most often single and in a subcapsular location, these lesions grossly appear as grey-white firm nodules that measure usually less than 1 cm in diameter.

Histologically, multiple small bile ducts are surrounded by a fibrous stroma with no evidence of increased cell turnover, mitoses or invasion of adjacent liver (25). The natural history and clinical significance of intrahepatic bile duct adenomas is unknown but rare malignant transformation has not been ruled out (26). Their main significance is their possible confusion with metastatic carcinoma during laparotomy, but pathological diagnosis following excisional biopsy is readily accomplished (23). A link between bile duct adenomas with GCTs elsewhere in the biliary tree has not been reported and whether there is a potential association remains open to speculation.

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