Spontaneous rupture of a nonparasitic hepatic cyst associated with peritonitis

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P SHIPLEY, B BAYLIS, N HERSHFIELD, R LUI, NCW WONG. Spontaneous rupture of a nonparasitic hepatic cyst associated with peritonitis. Can J Gastroenterol 1991;5(5):171-173. The first report of a nonparasitic cyst complicated by rupture and peritonitis is given. A 63-year-old female found to have a nonparasitic hepatic cyst was discharged from hospital when her symptoms of sharp intermittent pains in the right upper quadrant resolved spontaneously. Hours later, she was re-admitted with rupture and peritonitis. After hepatic cystojejunostomy (Roux-en-y) and T-tube placement in the common bile duct, the patient remains asymptomatic two years later. Key Words: Hepatic cyst, Nonparasitic, peritonitis, Rupture

Rupture spontanée d’un kyste hépatique d’origine non parasitaire et péritonite

RESUME: On rapporte le premier cas de kyste non parasitaire compliqué d’une rupture et d’une péritonite. Une patiente de 63 ans et chez qui on avait diagnostiqué un kyste hépatique avait pu regagner son domicile quand les douleurs vives et intermittentes qu’elle éprouvait à hauteur du quadrant supérieur-externe droit avaient spontanément cessé. Quelques heures plus tard, elle a été de nouveau hospitalisée souffrant d’une rupture de kyste et d’une péritonite. La patiente a subi une cystojejunostomie (opération de Roux en y) avec pose d’un drain de Kehr dans la voie biliaire principale. Elle est toujours asymptomatique deux ans plus tard.

IN MOST INCIDENCES NONPARASITIC hepatic cysts are uncomplicated, but the authors have recently encountered a case associated with spontaneous rupture leading to peritonitis. The occurrence of nonparasitic hepatic cysts is very low, clearly demonstrated by the results of a 15 year retrospective study at the Mayo Clinic in which 10,000 abdominal operations were surveyed and only 15 found to be due to a nonparasitic hepatic cyst (1,2). The finding of a nonparasitic cyst complicated by rupture and peritonitis is extremely rare, thus prompting the authors to summarize their experience in this report.

CASE PRESENTATION

A 63-year-old female was admitted to the Calgary Foothills General Hospital complaining of sharp intermittent pains in the right upper quadrant of the abdomen with radiation to the right shoulder and arm. Six months previously the patient had experienced episodes of recurrent fever, chills and fatigue. The only significant finding at
the time of admission was a tender globular mass located in the right upper quadrant of the abdomen, that moved with respiration. Auscultation of the mass failed to reveal a rub or bruit.

Laboratory investigations showed that white blood cell count (5.8x10^9/L), aspartate aminotransferase (25 U/L) and total bilirubin (11 µmol/L) were all within normal limits. Potential etiologies for hepatic cyst such as amebiasis and echinococcus were excluded by negative serology. The only positive finding was an ultrasound study showing a homogeneous solitary cyst (13x15 cm²) on the inferior surface of the right lobe of the liver. The location and size of this lesion was confirmed by computed tomography scan (Figure 1A). The course in hospital was unremarkable. The patient’s symptoms resolved spontaneously, and the patient was discharged.

Within hours of discharge, a sudden change in the characteristics of the pain prompted the patient to return to hospital. This time, the abdominal pain was severe and generalized. Physical examination on re-admission showed that the patient was in acute distress (blood pressure 140/80 mmHg, pulse 120 beats/min, and respiratory rate 24/min). Abdominal examination revealed a distended abdomen and signs of peritonitis. Significant laboratory findings included a white blood cell count of 15.6x10^9/L with 87% polymorphonuclear cells. A flat plate x-ray of the abdomen had the appearance of ground glass, suggesting the presence of fluid in the peritoneal cavity. Repeat computed tomography scan demonstrated that the cyst had ruptured and broken through Glissen’s capsule (Figure 1B). Medical management included intravenous fluids and an antibiotic, cefoxitin (2 g every 4 h).

After the patient’s condition had stabilized, she underwent hepatic cystojejunostomy (Roux-en-y) and T-tube placement in the common bile duct. Analysis of fluid from the cyst revealed both blood and bile, but cultures of the fluid were negative. An intraoperative T-tube cholangiogram failed to show communication of the cyst with the biliary tree. Histological reports indicated that the cyst was lined by low columnar epithelial cells of uniform thickness with no evidence of malignancy. The patient’s postoperative course was unremarkable. No complications or recurrence were encountered during a brief follow-up period (Figure 1C), and the patient remains asymptomatic two years following the procedure.

DISCUSSION
In the present report the authors have summarized the events leading up to the diagnosis and management of a nonparasitic cyst of the liver. The histological finding of low columnar epithelial cells indicates that the lesion was a congenital ductal cyst of the liver (2). Congenital hepatic cysts in the absence of polycystic kidney disease are very uncommon (2). The incidence of this disease is highest in elderly females (3). The single feature of this case presentation that sets it apart from others is the spontaneous rupture of the cyst leading to peritonitis.

Solitary congenital hepatic cysts are classified as being either parenchymal or ductal (4,5). Conservative management is usually indicated for asymptomatic solitary parenchymal or ductal cysts, but complications may occur. Parenchymal cysts arise from the tissue of mucous glands or from the teres ligamentum (6). Cysts of this nature give rise to symptoms of abdominal fullness and bile tract obstruction when they become filled with transudative fluid (3,7-10). In contrast, ductal cysts arise from the failure of provisional embryonic bile ducts to undergo involution (1,5,7,10). Despite the absence of communication between ductal cysts and major bile tracts (1,11), hemorrhage and bile leakage into the cyst are known complications (2,3,10).

The literature contains only three documented cases of nonparasitic hepatic cyst that ruptured, none of which were associated with peritonitis.
The authors have documented the present case as an example that goes against the general belief that rupture of a nonparasitic hepatic cyst is a benign condition. In light of this experience the authors suggest that more aggressive management may be indicated in certain cases. For example, cysts that contain transudates may be drained with transabdominal aspiration (4,13,14). External drainage is indicated for infected cysts. In contrast, bile- or blood-containing cysts can be safely drained by cystoscuticostomy. Because hepatic cyst rupture occurs most commonly in the elderly it is essential that early surgical intervention be weighed against the potential outcome described above, as this age group is least capable of tolerating severe illness.

In conclusion, the authors have reported a case of nonparasitic cyst of the liver that ruptured, leading to acute peritonitis. To their knowledge, this is the only reported case in which peritonitis has occurred following rupture of a nonparasitic hepatic cyst.

REFERENCES

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