Laparoscopic surgery for congenital biliary dilatation accompanied by accessory hepatic ducts: a report of seven cases

Yuebin Zhang, Jia Yu, Chen Qingjiang, Duote Cai, Zhigang Gao

Congenital biliary dilatation (CBD) is a congenital biliary tract anomaly characterized by expansion of the common bile duct and is thus known clinically as a choledochal cyst. The form of CBD varies, and the commonly used classification was described by Todani et al. Pancreaticobiliary maljunction is reported to be present in 30%-96% of patients with CBD. The main aims of laparoscopic surgery include resection of the dilated biliary ducts, gallbladder, pancreatic-bile diversion, and biliary tract reconstruction. Accessory hepatic ducts (AHDs) are extrasegmental bile ducts that often drain a part of the liver and can join the biliary system at any position, such as the common bile duct, common hepatic duct (CHD), right hepatic duct or the gallbladder. AHDs are seldom seen on preoperative imaging, but they increase the difficulty of surgery and the risk of postoperative biliary fistula. A retrospective analysis of treatment outcome in 7 of 542 patients with AHDs who received surgical treatment in our center from January 2012 to August 2020 was conducted.

A total of 542 patients received CBD surgery in the Department of General Surgery, Zhejiang University Children's Hospital from January 2012 to August 2020. Most of the patients presented with typical choledochal dilatation during surgery and underwent classic cholecystectomy with dilated biliary duct and Roux-en-Y anastomosis of the hepatic duct and jejunum. Of these 542 cases, 7 patients had anatomic variations of AHDs, and choledochojejunostomy was performed differently. Of these seven patients, two were male and five were female, with an average age of 23.5 months, ranging from 2 to 48 months. Only one case was identified by preoperative imaging using magnetic resonance cholangiopancreatography (MRCP) as having AHDs (figure 1), and the rest were found intraoperatively. Of these seven cases, two were found to have choledochal cysts during prenatal examination, five cases were found to be due to abdominal pain in hospital examination, and two of them were found to be abnormal liver function with increased direct bilirubin of 61.6 µmol/L and 35.3 µmol/L, alanine aminotransferase (ALT) of 119 U/L and 78 U/L, and γ-glutamyltransferase of 1278 U/L and 496 U/L. Routine preoperative examination included routine blood tests, C reactive protein level, liver function, coagulation function, orthotopic chest radiography, electrocardiogram (ECG), abdominal B-mode ultrasound and MRCP.

The patients were placed in the supine position with the head high and feet low. A 5 mm 30° laparoscope was used. Four punctures for 1–2 5 mm trocars and the rest were 3 mm trocars were located in the left upper abdomen, right upper abdomen, navel and right lower abdomen, respectively. The anatomical separation of the extrahepatic biliary tract and treatment of the jejunal segment were basically the same as those for typical CBD, the jejuno-jejunostomy was extracorporeal, the proximal intestine and biliary branches were retained at 15–25 cm. The difference was mainly in the hepaticeojunostomy during surgery. All the operations were performed under the 'four-hole method' laparoscopy, and there were no cases of conversion to laparotomy. The average operation time was 157 min (range 145–210 min) compared with 142 min without AHD, and intraoperative bleeding was less than 10 mL. AHD was found in one case by preoperative magnetic resonance imaging (MRI) examination, abnormal bile duct was found in four cases when the neck of gallbladder was dissected, and AHD was found in the other two cases when bile exudation was found on the wound surface during irrigation. The
The diameter of AHD was obtained by comparing with the tip of the laparoscopic forceps, and the diameter of the electrocoagulation hook was considered to be 1 mm (figure 2).

The seven cases of CBD with AHD variation were treated differently according to the size and location of the AHD and were divided into three surgical methods as shown in table 1. Method 1: after intermittent suture of the AHD and CHD, a fusiform choledochal-jejunum anastomosis was performed (three cases in total), which is shown in figure 3. Method 2: the AHD and CHD were anastomosed with the jejunum, respectively (two anastomotic stoma), in two cases (figure 4). Method 3: ligation of the AHD (with diameter less than 2 mm) in two cases (figure 5).

CBD is a common congenital biliary tract anomaly. The preferred radical treatment is biliary tract resection with biliary tract Roux-en-Y reconstruction. In 1995, Farello et al first reported laparoscopic surgery for CBD. In 2002, Li et al first reported laparoscopic treatment of CBD in China. In children with CBD, the presence of a variant bile duct and AHD also may affect the surgical procedure. The rate of biliary tract variations is high, and the variation forms are complex and changeable. The classic normal biliary tract structure accounts for only 57.6%, and the variation rate can be as high as 42.4%. AHDs are variations of the biliary tract that drain the bile ducts of one lobe or segment of the liver, join into the gallbladder, common bile duct, CHD, and right hepatic duct. It is reported that the incidence of AHDs is 1.0%-31.4%. However, the proportion of AHDs encountered during CBD surgery is not high, and the diameters of the AHDs are usually small. Most of them cannot be clearly defined by preoperative MRCP and intraoperative angiography. In most cases, bile extravasation is often detected by careful examination during the operation. We found seven cases complicated by AHDs during surgery, and the incidence was approximately 1.3% (7 of 542 cases). Children’s AHDs are often found to be only 1-2 mm in diameter, and both preoperative ultrasound and MRCP are often unable to clearly show the shape of AHDs. We do not regularly perform preoperative ERCP imaging in children with CBD, and intraoperative biliary angiography cannot clearly identify AHDs because the image is overlapped by the gallbladder or choledochal cyst. The presence of AHDs is identified by the presence of small ducts or bile leakage during surgery (figure 4). The Japanese scholar Takahashi summarized five CBD cases of AHDs: only one case was found during intraoperative imaging, one case was found by repeated endoscopic retrograde cholangiopancreatography (ERCP) imaging, and the rest were found by intraoperative or postoperative biliary fistula. It has been reported in the literature that more cases of AHD were found due to bile leakage during surgery or due to the postoperative complication of biliary fistula. In this group of seven cases, preoperative imaging with MRCP showed the presence of AHD in only one case, and in most of the children the presence of AHD before surgery could not be predicted. In four cases, small ducts were found during the operation, which were confirmed to be AHDs after careful dissection. In two cases, bile leakage was found during rinse examination of wound surface after dissection, and AHD injury was found on subsequent examination; thus, careful assessment of anatomy during the operation is particularly important. The junctions of the AHDs in these seven children were located in the gallbladder or in the neck of the gallbladder duct. During the operation, the anatomy of this site should be monitored carefully to distinguish the aberrant blood vessels from the AHD, and it is also necessary to wash the wound surface.
during the operation to avoid the occurrence of postoperative bile leakage. If the diameter of the AHD is more than 2 mm, it is feasible to separate the AHD by careful assessment of anatomy during the operation when there is no chronic inflammation of the biliary tract and local adhesions are not serious. If the AHD is too small (e.g., diameter <2 mm) or if the anatomical region is not clear because of repeated infection of the cyst, it is difficult to avoid injury to the AHD. The AHD has the function of bile drainage, and anastomosis should be carried out as far as possible if injury of the AHD is found during the operation. Wen Zhe proposed that biliary tract reconstruction is recommended for AHD over 2 mm.12 In our center, three of seven cases whose AHDs (approximately 2 mm in diameter) were around the CHD were anastomosed to the branch of the jejunal biliary tract with a spindle shape after intermittent suture of the AHD and the lateral wall of the CHD to ensure bile drainage. Two of seven cases whose AHD diameter was 2 mm but also a certain distance from the CHD could not be combined together. Therefore, two choledochojejunostomies were performed separately. In the remaining two cases whose AHD diameter was less than 2 mm, which was the non-anastomotic type, simple ligation was performed. After the above treatment, all patients were followed up for 2–23 months. None of the patients had abnormal liver function or elevated bilirubin, and B-mode ultrasound examination showed no dilatation of the biliary tract inside and outside the liver. It is safe and feasible to ligate the AHD with a diameter less than 2 mm. Although anastomosis of the AHD with a diameter greater than 2 mm is difficult and requires an experienced surgeon, postoperative complications, such as increased bilirubin, bile fistula and biliary stricture are rare; thus, anastomosis is recommended. If the AHD is close to the hepatic duct, a spindle anastomosis after tenseless combination can reduce the difficulty of anastomosis, which is a safe and effective surgical method.

Postoperative recovery was uneventful. No postoperative bleeding, biliary fistula, intestinal fistula, biliary obstruction or ileus complications occurred during hospitalization. The average postoperative hospital stay was 10 days and ranged from 8 to 14 days. For anastomotic stenosis occurring months to years after surgery, long-term follow-up is maintained. The postoperative follow-up period ranged from 2 months to 23 months, and the follow-up indices measured included serum bilirubin, ALT, routine blood tests, C reactive protein and ultrasonography. For the two cases with abnormal liver function before surgery, with direct bilirubin of 61.6 µmol/L, ALT of 119 U/L and γ-glutamyltransferase of 1278 U/L, these values returned to normal 19 days after surgery in case 1.

**Table 1** Seven cases of CBD associated with AHD

<table>
<thead>
<tr>
<th>Case</th>
<th>Sex</th>
<th>Diameter (mm)</th>
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<td>7</td>
<td>F</td>
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</table>

AHD, accessory hepatic duct; CBD, congenital biliary dilatation; F, female; M, male; MRCP, magnetic resonance cholangiopancreatography; OP, operation.

**Figure 3** Anastomosed to the branch of the jejunal biliary tract with a spindle shape after intermittent suture of the accessory hepatic duct (AHD) and the lateral wall of the common hepatic duct (CHD).

**Figure 4** AHD and CHD were anastomosed with the jejunum, respectively (two anastomotic stomas). AHD, accessory hepatic duct; CHD, common hepatic duct.
Direct bilirubin of 35.3 µmol/L, ALT of 78 U/L and γ-glutamyltransferase of 496 U/L returned to normal 13 days after surgery in case 2. The seven patients with CBD and AHDs did not have an abnormal increase in bilirubin, and biliary ultrasonography showed no obvious dilatation of the biliary tract inside and outside the liver 2–23 months after surgery.

In summary, biliary ducts of patients with CBD have varied dilated locations and forms and are often associated with variable biliary ducts and blood vessels. For the treatment of AHD, it is necessary to dissect and identify the abnormal vessels and AHDs carefully during the operation, and it is also necessary to wash the wound after the dissection. If the diameter of the AHD is larger than 2 mm, choledochojejunostomy should be performed. The surgical methods included two independent choledochojjunostomies when the AHD is far from the CHD, and one spindle anastomosis when the AHD and CHD are close.

Figure 5  Ligation of the AHD. AHD, accessory hepatic duct; CHD, common hepatic duct.

REFERENCES

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