Management of Acute Ascending Cholangitis with Septic Shock on Top of Altered Anatomy after Old Road Traffic Accident: A Case Report

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Abstract

Introduction: Acute cholangitis is a clinical syndrome characterized by fever, jaundice, and abdominal pain that develops because of stasis and infection in the biliary tract. The most frequent causes of biliary obstruction are biliary calculi, benign biliary stricture, and malignancy.

Case presentation: Here we report a case presented with septic shock and acute ascending cholangitis on top of an altered anatomy in the form of disruption of the head and Wirsung duct of the pancreas, complete avulsion of the CBD at the ampulla of vater and deformed duodenal bulb after an old road traffic accident sustained thirty-five years back. Initial resuscitation, and antibiotic therapy with urgent decompression have been done. Definitive surgery has been performed in the form of dismantling of prior cholecystojejunostomy, exploration and resection of common bile duct with cholecystectomy and Roux-en-Y hepaticojunostomy with adhesiolysis.

Conclusions: Definitive surgery remained the mainstay of treatment for extrahepatic biliary and pancreatic trauma to prevent future life-threatening complications. MDT approach is the optional choice for dealing with critically ill patients. Utilization of the full capabilities of endoscopic and interventional radiology services deemed mandatory for urgent and timely management.

Keywords: Acute cholangitis; Syndrome; Radiograph; Duodenal bulb

Introduction

Acute cholangitis is a clinical syndrome characterized by fever, jaundice, and abdominal pain that develops because of stasis and infection in the biliary tract. It is also referred to as ascending cholangitis. Cholangitis was first described by Charcot as a serious and life-threatening illness; however, it is now recognized that the severity can range from mild to life-threatening [1]. The most frequent causes of biliary obstruction in patients with acute cholangitis without bile duct stents are biliary calculi (28% to 70%), benign biliary stricture (5% to 28%), and malignancy (10% to 57%) [2]. Acute cholangitis can also occur following endoscopic retrograde cholangiopancreatography (ERCP) (0.5% to 1.7%), particularly therapeutic ERCP following stent placement, or postoperatively due to bile duct injury, or a stricture of biliary-enteric anastomosis. Rarely, the distal common bile duct may be obstructed by food, stones, or debris in patients with a biliary-enteric anastomosis (Sump syndrome) [3-5]. Here we report a case presented with septic shock and acute ascending cholangitis on top of an altered anatomy after an old road traffic accident sustained thirty-five years back.

Case Presentation

A 63-year-old male, who has a known case of diabetes mellitus on oral hypoglycemic presented to ED with an epigastric pain of 2 days duration with a pain score of 10. He complained of nausea, vomiting, anorexia and chills as well. He has no known allergy; however, he has a significant surgical history of blunt abdominal trauma sustained thirty-five years ago. Such injuries which were vague and not known in a detailed manner on presentation as the patient had been managed in another country at the time. At presentation he was in shock and feverish; BP: 80/40 -HR: 106-Tem: 40.1°C –RR: 20/min -SO₂ 98% on 2L via nasal cannula. On examination, clinical jaundice was noticed, in addition to a median laparotomy scar. His abdominal examination showed moderate tenderness in the epigastric region. On presentation, he had an open bowel. The patient was managed as per ACLS protocol with fluid resuscitation and noradrenaline 0.5 mcg/kg/min started. RBS checked and revealed 17.6 mmol/L, so insulin infusion started. Laboratory results were as follows; Na: 122-K: 4.09-Crea: 118-Urea: 8.61-WBC: 9.31-Neut 94%-Hb: 13.1 PLT: 168-AST: 219-ALP: 394-ALT: 227-GGT: 557-total bilirubin: 86-Direct bilirubin: 80-Amylase: 19.09-Lipase: 56 -CRP: 143. H1N1 -ve. VBG: PH: 7.22, PCO₂: 46, HCO₃: 18, Lac: 8.01. He had a sinus rhythm on ECG. Bedside ECHO was normal with preserved LVEF 55%.

Chest radiograph showed accentuated lung markings with multiple short air-fluid levels at the right side of the abdomen. On the abdominal radiograph, multiple faint radio-opaque shadows were noticed at the right hypochondrium, query GB stones. The patient had been admitted to ICU and sepsis protocol was applied. Furthermore, antibiotics were administered (Meropenem-Ciprofloxacin-Vancomycin) in addition to fluconazole. Abdominal ultrasound was done (Figures 1A and 1B) revealing dilated CBD of 9.5 mm diameter with single stone seen at its proximal part measuring 10 x 7 mm. Dilated IHBRs were seen at both hepatic lobes. The gallbladder was contracted over stones with a thickened edematous wall. Urgent decompression by ERCP was deemed necessary. Unfortunately, cannulation failed even with precut needle knife.

In the view of persistent tachycardia and high inotropic support, the patient was intubated electively for PTC, where cholangiogram was done which revealed IHBRD with dilated CBD and filling defects seen within likely of calcular nature. No obvious communication of distal end of CBD with the duodenum but only localized fixed opacified area

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One day later, Echocardiogram was done giving the picture of type II myocardial injury due to underlying sepsis, with impaired systolic function-EF 30%, markedly hypokinetic mid-lateral and mid to basal infero-lateral segments with hypokinetic remaining segments. Grade II LV diastolic dysfunction. Trivial MR. Mild to moderate TR with estimated systolic PAP of 35 mmHg. Cardiac profile showed CK: 340-CMB: 9.53-Troponin-I: 11.1. The patient’s situation had become clearer with ascending cholangitis, obstructive jaundice, septic shock, AKI, type 2 myocardial injury and deranged coagulation. Laboratory results were: WBC: 24.32-CRP: 202-GGT: 249-ALP: 172-T: Bil: 72.1-D. Bil. 57.64-Ure: 13.7-Creat: 177.6-ALT: 97-AST: 101-INR: 2.02-APTT: 48.4- APTT ratio: 1.86-PT: 23.2. Blood culture grew Escherichia coli.

Drainage from PTC catheter was nil and because of the deteriorated critical situation of the patient, decision had been taken to upsize the PTC catheter for decompressing his biliary system via 12 F catheter (Figures 3A and 3B). On the third day of admission, the newly inserted PTC catheter drained 110 cc and abdominal ultrasound was done (Figures 4A-4D) which showed PTC drainage tube inserted. Significant improvement regarding the previously seen dilatation of IHBRs and CBD was seen. A newly seen mild amount of free fluid at right iliac fossa was newly seen.

One day later, PTC drained 190 cc clear bile, however, LFT was noted trending up with T: Bil: 108.3-D: Bil. 80.67. WBC also jumped to 32.12 with 90% neutrophilia. RFT manifested a significant rise; urea measured 23.7 and Create was 224.73. Moreover, the patient had gaseous distension of the abdomen that was confirmed by abdominal radiograph and hypoactive bowel sound with last bowel motion on the day of admission. MDT meeting gathered which in turn agreed to do a US abdomen and compare it with the previous one. CT abdomen with contrast would have been required if any significant abnormal finding in the US or any deterioration of the patient’s condition had ensued. Results of US (Figures 5A-5E) were; more improvement regarding the previously seen dilatation of IHBRs and CBD with a change in the location, but with the same amount of free fluid seen at left iliac fossa due to change patient position. The gallbladder became more distended and shows large stone inside measuring 10 x 8 mm with a significant decrease in gallbladder wall thickness (from 12 mm to 6 mm). A newly seen rim of free fluid at peri-hepatic recess was noticed.

Over the next 7 days, the patient’s condition had been dramatically improved. He was extubated, hemodynamically stable without inotropic support, shifted to the ward, the urinary catheter was removed, started oral feeding and opened his bowel. Laboratory findings had correlated well with the ameliorated clinical condition where, Urea: 5.16-Creat: 33.08-WBC: 7.9-Nuet 73.8%-CRP: 41.7-T: Bil. 59.6-D. Bil: 41.2-GGT: 281-ALP: 239. Cholangiogram was done at the time (Figures 6A and 6B) after removal of the previously inserted external biliary drainage catheter over guidewire-which showed dilated tortuous cystic duct with opacification of the proximal CBD distal to the cystic duct starting from confluence site with two distal CBD large filling defects likely of CBD stones, opacification of gallbladder as well as the gallbladder fundus-jejunum anastomotic site with a detectable multiple gall bladder stones of variable size. Advancement of external-internal 8.5F biliary drainage catheter was done with its internal end within the jejunal loop.

Over another week in the ward the patient condition had remained stable and planning for definitive surgical management was commenced, an official report of the old trauma was obtained which documented the following injuries sustained on this RTA, huge retroperitoneal bile stained haematoma, disruption of the head and Wirsung duct of pancreas, complete avulsion of the CBD at the ampulla of Vater and two duodenal perforations posteriorly and medially. Such injuries were managed by closure of duodenal perforations, T-tube splint of the CBD, cholecystojejunostomy and sump drain insertion near the head of the pancreas. Postoperative investigations after the acute clinical course of his old trauma revealed that; on HIDA scan, bile enter the GIT through the cholecystojejunostomy. On ERCP contrast fails to enter either CBD or pancreatic duct, double contrast study showed deformed duodenal bulb. At the time since the patient was not complaining of steatorrhea nor jaundice he was advised to continue follow up with a possible need to convert his cholecystojejunostomy to choledochojejunostomy after further work up.

Taken into account this complicated anatomy, work up for definitive management started with CT with oral and IV contrast (Figures 7A-7C) which confirmed the following: sizable subcapsular collection along the right lateral and cranial aspects of the liver, with a maximal thickness of 4 cm. In addition, internal/external biliary drainage catheter is noted coursing through the collection, liver, gallbladder and extending to the jejunal loops. There is also air within the lumen of the gallbladder and small amount of fluid within the Morison’s pouch. The pancreas is not visualized in keeping with near complete lipomatosis and suturets were noted along the wall of the hepatic flexure of the colon.

Despite the continuing clinical improvement, the discovery of perihepatic collection mandated prompt intervention where, a US guided 12F drainage catheter was inserted (Figures 8A and 8B) within the perihepatic collection yielding clear bile, which was removed after five days when the drainage was about nil. Echocardiography was repeated showing normal left ventricular cavity size with preserved global systolic function (EF 55%, previously 35%) with no regional wall motion abnormality. The patient was discharged home with the PTC drain, in a good condition with a plan for definitive surgery after maximal optimization of his general condition.

Ten days later the patient came to hepatobiliary surgery clinic for assessment, the abdomen was soft and lax with midline laparotomy scar and 2 previous drain scars. PTC drain was connected and functioning. Management Plan was explained to the patient and his family, clarifying the need to have a definitive procedure to avoid future cholangitis and sepsis. In addition, the complexity and possible complications of the operative procedure (cholecystectomy plus excision of the previous cholecystojejunostomy with reconstruction of hepaticojejunostomy) were emphasized with cons and pros vs. endoscopic interventions.

An MDT meeting with radiology, GI and HB surgery was arranged. Referral to cardiology for assessment of fitness for surgery, a dietitian for high protein low carbohydrates diet and endocrinology for blood sugar control were done. Eventually, the patient was sent for preoperative anaesthesia consultation and booked for surgery after full optimization of his general condition and preoperative cholangiogram (Figure 9), which excluded any stricture or abnormal dilatation of intrahepatic biliary radicles, hepatic ducts or proximal CBD and confirmed the obstruction of CBD and dilatation and tortuosity of the cystic duct. The contrast flew from the gallbladder to the jejunum via cholecystojejunostomy.

The patient was admitted electively for exploratory laparotomy with dismantling of prior cholecystojejunostomy, exploration and resection of the common bile duct with cholecystectomy and Roux-en-Y hepaticojejunostomy with adhesiolysis.
Figure 1: (A) CBD dilated with a stone within. (B) Dilated IHBR.

Figure 2: (A) Cholangiogram showing IHBRD, dilated CBD and filling defects seen within, with no communication of distal end of CBD with the duodenum. (B) 8.5 F external biliary drainage catheter in place.

Figure 3: (A) Upsizing of the previously inserted 8.5. (B) F external biliary drainage was done by bigger one-12 F.

Figure 4: (A) PTC drainage tube inserted. (B) Free fluid at the right iliac fossa. (C) Significant improvement regarding the previously seen dilatation of IHBRs and CBD. (D) Contracted gallbladder with a thickened edematous wall with a stone within its lumen.

Figure 5: (A) Improvement of dilatation of IHBRs and CBD. (B) The gallbladder becomes more distended and shows large stone inside measures 10 × 8 mm. (C) Rim of free fluid at peri-hepatic recess. (D) Drainage tube in place. (E) Change in the location but with the same amount of free fluid seen now at left iliac fossa.

Figure 6: (A) Dilated tortuous cystic duct with opacification of the proximal CBD distal to the cystic duct starting from confluence site with two distal CBD large filling defects. (B) Opacification of gallbladder as well as the gallbladder fundus- jejunal anastomotic site.

Figure 7: (A) Sizable subcapsular collection along the right lateral and cranial aspects of the liver. (B) Internal/external biliary drainage catheter is noted. (C) There is air within the lumen of the gallbladder with evidence of cholecystojejunostomy.
Discussion

Obstruction of the biliary ducts and presence of a superposing bacterial infection are common features in cholangitis. Not every biliary obstruction is associated with cholangitis but there is surely a biliary obstruction in every cholangitis case [6-9]. Biliary obstruction is caused by choledochothiasis mostly. Moreover, malignancy, benign strictures, and interventions to the biliary ducts may be the cause of biliary obstruction. Other mechanisms include extrinsic compression of the bile duct due to a duodenal periampullary diverticulum (Lemmel syndrome), inflammation secondary to acute pancreatitis, or an impacted stone in the cystic duct or neck of the gallbladder (Mirizzi syndrome). Other intrinsic causes of biliary obstruction include blood clots, and parasitic infections (mainly liver flukes and the roundworm Ascaris). Retained worm fragments can serve as a nidus for biliary stones and cause recurrent pyogenic cholangitis. Many other rare causes of cholangitis are reported in the literature, like recurrent cholangitis without stones with previous papillotomy, a picture suggestive of a causal relationship between heavy lifting and cholangitis, where increased intra-abdominal pressure can cause reflux into the common bile duct in the presence of a papillotomy [10]. A similar aetiology of recurrent cholangitis was described in patients who had undergone biliary bypass or pancreateoduodenectomy [11]. Another case report revealed another rare mechanism, where the condition was attributed to food impaction through a choledochoduodenal fistula leading to obstructive jaundice with ascending cholangitis progressing to sepsis [12]. External compression of the biliary tree from a ruptured renal angiomyolipoma (AML) of the right kidney and mucinous cystic neoplasm of the liver with extrahepatic growth have been reported as rare causes of ascending cholangitis [13,14]. A more unusual case of ascending cholangitis has been reported due to fractured ventriculo-gall bladder shunt fragments causing mechanical obstruction [15].

Common to all forms of biliary obstruction is elevated choledochal pressure which is deemed the inciting factor for the inflammatory process and sepsis. The biliary system works according to a pressure gradient where the pressure of the hepatic bile secretion is 120-150 cm H₂O and the pressure in extrahepatic bile ducts is 100-150 cm H₂O. So normally, bile secretion occurs according to these pressure values and bile fills into the gallbladder with a pressure of 12-18 cm H₂O. Peristaltic contraction-relaxation of the sphincter of Oddi is the most important factor in the regulation of this pressure. Bile secretion from the liver is inhibited if the pressure exceeds 300 cm H₂O. If the choledochal pressure exceeds 25 cm H₂O, hepatic defense mechanisms against infection become useless, consequently, in 25-40% of the cases, the associated infection spreads into the intrahepatic canaliculi, and cholangio-venous reflux ensues, followed by the access to the hepatic veins and lymphatics, resulting in bacteriemia. Sepsis is common in the case of suppurative infections [16,17]. Isolated organisms are E. coli (27%), Klebsiella (16%), Enterococcus (15%), Streptococcus (8%), Enterobacter (7%), and Pseudomonas aeruginosa (7%) [18,19].

In our patient with altered anatomy multifactorial pathogenesis played a role to initiate such stormy clinical course. It is well known that risk factors for stone formation are bile duct dilatation, biliary stricture, and angulation of the common bile duct due to anatomical abnormalities [20,21]. The obstructed CBD, tortuous cystic duct and distorted duodenal bulb contributed to the formation of his CBD and gallbladder stones which in-turn led to the obstruction and creation of a vicious circle. Although bacterial colonization of the small intestine is not uncommon, it is rarely associated with widespread sepsis. Colonization itself is often a consequence of the failure of intestinal clearance of bacteria which may be due to dysmotility or altered anatomy. The prevalence of bacterial overgrowth varies in association with the predisposing medical disease or surgically modified anatomy, and whether patients studied are symptomatic [22-25]. Again, the altered anatomy with direct continuity of the patient biliary system with the jejunum through the previously formed cholecystojejunostomy provided a portal of entry through which E. coli gained access to his biliary system to induce his cholangitis and septic shock, where it was isolated from his blood culture.

Common bile duct (CBD) injuries from blunt abdominal trauma are rare. In fact, extrahepatic biliary tract injuries occur in 3% to 5% of all abdominal trauma victims, with 85% resulting from penetrating wounds. Of the remaining 15%, resulting from blunt trauma, the clear majority, 85%, involve the gallbladder alone [26-28]. CBD injury occurs frequently at three areas of relative fixation of the biliary tract [29]:

1. The origin of the left hepatic duct,
2. The bifurcation of the hepatic ducts,
3. The pancreatocoduodenal junction.

Different mechanisms, even in combination, may produce rupture of the common bile duct: Compression of the ductal system against the vertebral column [30], sudden increase of intraluminal pressure in the gallbladder with a short and permeable cystic duct [31], and a “shearing force” producing avulsion of the common duct at its fixed part at the junction with the pancreas [32]. Regarding pancreatic injury, it is also rare but severe complications occurring in 2-5% of blunt abdominal trauma [33,34] and are usually associated with injuries of other intra-
abdominal organs. Isolated pancreatic rupture is quite rare due to its location in a relatively protected retroperitoneal area of the abdominal cavity [35,36]. In the acute setting, clinical symptoms are often initially subtle, leading to delay in diagnosis, increase in morbidity and mortality of up to 60% [37]. The common complications of pancreatic injuries are a pancreatic fistula, intra-abdominal abscess, pancreatitis, pseudocyst and sepsis [38].

The treatment options for an extrabiliary biliary leak have broadened. Until recently, such injuries usually mandated surgical repair utilizing debridement and closure with or without T-tube; patch closure using gallbladder, cystic duct, vein, serosa or jejunum; biliary-enteric anastomosis using duodenum or jejunum; or liglation and drainage with plans for subsequent enteric diversion [39]. For pancreatic injuries, the integrity of the main pancreatic duct (MPD) is the most important determinant of prognosis in these patients, with disruption to the MPD an indication for laparotomy [40,41].

Pancreatic injuries are classified into five grades according to the Pancreas Injury Scale published in 1990 by the American Association for the Surgery of Trauma [42]. Traditionally, patients with injuries to the MPD (grade III, IV, V) require laparotomy. However, a trial of non-operative treatment may be considered even in more severe injuries such as isolated grade IV injuries if the patient is haemodynamically stable. Endoscopic stenting and drainage is an attractive minimally invasive therapeutic procedure for haemodynamically stable patients, and it may obviate the need for surgery [43-46].

From the aforementioned, we can get an integrated image of the management of the acute extrabiliary biliary system and pancreatic injuries. However, for chronic deformities and altered anatomy, the literature has scanty resources. The presentation of our patient with injuries to the extrahepatic biliary system and pancreas illustrates the need to consider the patient as a whole with many organ systems involved. The patient was admitted for definitive surgery.

Operative details

Midline incision with J extension to the right side from inferior aspect was performed where extensive adhesions were encountered from previous surgeries. Careful adhesiolysis was done to separate the liver from the colon, small bowel and stomach. Prior choledochojunostomy was identified and dismantled with Endo GIA stapler and the gallbladder was dissected down to the confluence with the CBD. Then the CBD and the CHD were clearly identified. The CBD was transected distally, and its stump was closed in a continuous fashion after ensuring that, no stones were left in the remnant. The CHD was then divided with the removal of the gallbladder and the CBD. The jejunum was divided with endo GIA at the same site as the previous choledochojunostomy. Jejunoojejunostomy was created with the alimentary limb being anastomosed to the biliary limb with endo GIA and overrunning sutures. Finally, hepaticojejunostomy was created with Maxon 5 - 0 sutures in an interrupted fashion. The mesenteric defect was then closed to prevent a possible internal herniation. No bile leak was found post anastomosis. The peritoneal cavity was thoroughly washed with warm saline. Two 15 Fr JP drains were inserted, one at the site of biliary anastomosis and the other in the peritoneal cavity. Wound infusion catheters were placed for postoperative analgesia using bupivacaine. Hemostasis was achieved, and the abdominal wall was closed in 2 layers with looped No 1 PDS, while the skin was closed with staples.

The patient went through an uneventful smooth postoperative course with no single complication and came for follow up after discharge to our clinic, symptom-free, where skin clips were removed, and the patient reassured that his surgery was successful.

Conclusions

- Definitive surgery remained the mainstay of treatment for extrabiliary biliary and pancreatic trauma to prevent future life-threatening complications.
- MDT approach is the optimal choice for dealing with critically ill patients.
- Utilization of the full capabilities of endoscopic and interventional radiology services was mandatory for urgent and timely management.

References


