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Portal biliopathy
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Abstract
Portal biliopathy (PB) is a rare disorder, which mostly presents as sub-clinically. It occurs most commonly due to idiopathic extrahepatic portal vein obstruction. We present three cases having features of portal biliopathy secondary to portal hypertension. Our first case did not have a prior history of chronic liver disease while next two patients had previous history of chronic liver disease resulting in portal hypertension. Cavernous transformation of the portal vein due to extrahepatic portal vein obstruction is not infrequent but biliary obstruction in association with this disorder is distinctly uncommon. Proper case management is very important as prolonged biliary duct obstruction can lead to the development of ascending cholangitis or later on secondary biliary cirrhosis.

Keywords: Portal biliopathy, Portal vein obstruction, Portal hypertension.

Introduction
Portal biliopathy can be defined as biliary ductal and gall bladder wall abnormalities secondary to portal hypertension, more commonly due to portal vein thrombosis. In 30% cases of portal biliopathy, extrahepatic portal hypertension is the predisposing factor, in as many as half of the patients there is no predisposing cause. The changes in biliary and portal channels include dilated intra and extrahepatic biliary ducts, varices formation around porta hepatis due to portal hypertension caused by the portal vein obstruction. This can rarely result in cholelithiasis and choledocholithiasis formation secondary to biliary stasis.

The major presenting features of portal biliopathy are variceal bleeding, ascites, splenomegaly and encephalopathy. Jaundice is rarely seen in portal hypertension except in patients with history of underlying chronic liver disease. We present three cases having features of portal biliopathy secondary to portal hypertension. Our first case did not have prior history of chronic liver disease while the next two patients had a previous history of chronic liver disease resulting in portal hypertension.
Case-1:

A 30-year-old young gentleman presented with history of fever with chills and dry cough for the past five days. He also had epigastric burning sensation for which he took Ranitidine and oral antacids. Patient also contacted chest tuberculosis four years back for which he was treated for one year. The physical examination showed that his vitals were normal. He had jaundice however, and no anaemia. Systemic examination was also normal with soft, non-tender abdomen and no visceromegaly. Laboratory examinations included serum bilirubin of 4.4 mg/dl and raised alkaline phosphatase of 478 IU/L. Haemoglobin was 13.2 mg/dl and WBC of 5.7x10E9 with 66% neutrophils. Viral profiles including Hepatitis A, B, C, D and E were negative.

Ultrasound and CT scan were performed at the Department of Radiology, The Aga Khan University Hospital as diagnostic procedures. Extensive intrahepatic biliary dilatation was noted with numerous intrahepatic biliary duct calculi and sludge on ultrasound examination (Figure-1-a). However Common bile duct (CBD) could not be evaluated because of extensive peripancreatic and porta-hepatis varices (Figure-1-b). Varices were also seen at splenic hilum. No abdominal lymphadenopathy or ascites were noted.

Further workup with CT scan redemonstrated intrahepatic biliary dilatation with non-visualization of extra hepatic part of CBD. Low-density material was observed in intrahepatic ducts suggestive of sludge/calculi. Portal vein was not visualized separately. Marked collateral vessels were seen in the region of gastro hepatic ligaments extending into the porta-hepatis down to the level of renal hilum. However; there were no focal lesions or changes suggestive of chronic liver disease. Percutaneous transhepatic decompression of biliary system was performed to relieve the patients signs and symptoms due to biliary obstruction. Therefore, based on clinical and imaging findings, diagnosis of portal biliopathy was made by the radiologist. Further follow-up of the patient was not available as the patient was lost to follow-up.

Case-2:

A 19-year-old female, known case of hepatitis C, Protein C and S deficiency, presented with bloody vomiting and black colour stools for 1 day in Emergency Department. Vitals showed tachycardia with hypotension. Physical examination was unremarkable. Her liver function test was performed which was within normal limits. Direct Bilirubin was 0.5 mg/dl while alkaline phosphatase was 98 IU/L.
Follow-up alkaline phosphatase values were 144 and 359 IU/L respectively.

After initial assessment and management, endoscopy was done and band ligation was performed. Subsequently, ultrasound and CT abdomen were carried out which showed moderate intrahepatic biliary dilatation, gallstones, thrombosed portal and superior mesenteric veins, aneurysmal dilatation of the portal vein along with multiple dilated collateral seen at the porta hepatis, which were compressing the CBD at its mid level (Figure-2). Distal CBD at the head of the pancreas showed normal caliber, thus based on radiological features, diagnosis of Portal Biliopathy was made.

Furthermore, patient was followed in outpatient clinics. Multiple admissions were made because of recurrent cholangitis. Elective cholecystectomy followed by biliary stent placement was done via ERCP and patient now showed improvement in her symptoms.

Case-3:

This is another case of a 30 year old young male who was a known case of chronic liver disease and presented with history of abdominal pain in the emergency room. His initial lab investigations showed raised total bilirubin of 1.4 and direct bilirubin of 0.2. Alkalaine phosphatase was in normal range of 122 IU/L. Hepatitis B surface antigen and HCV Antibody were negative. He also had a history of upper gastrointestinal bleed six months back. Endoscopy was performed at that time which showed Grade IV varices for which band ligation was done outside of our hospital. Ultrasound at that time showed features of chronic liver disease with cavernous transformation.

CT scan was ordered as a workup procedure which showed cavernous transformation of portal vein with multiple collaterals at porta hepatitis and peripancreatic region resulting in mild intrahepatic biliary dilatation secondary to compression of distal common bile duct by these varices (Figure-3-a,b). Features of chronic bile duct dilatation and stent placement.

Figure-2 (b): ERCP showing s narrowing of CBD with intrahepatic biliary dilatation and stent placement.

Figure-3 (a,b): CT Scan: Multiple collaterals at porta hepatitis with intrahepatic biliary duct dilatation.

Case-3:

Discussion

Portal biliopathy is not the only cause of portal hypertension, there can be other intrahepatic and extrahepatic causes. While the extrahepatic causes include congenital portal vein obstruction, umbilical sepsis, trauma, hypercoagulation state like Budd-Chiari Syndrome and malignant occlusion, intrahepatic causes can be cirrhosis and primary biliary cirrhosis.1,4

The extrahepatic portal vein thrombosis causes formation of collateral channels mainly around porta hepatitis,
gastro hepatic ligament and splenic vein. These collaterals sometimes become large enough to cause compression and obstruction of common bile duct extrahepatically or intrahepatically. This occurs because bile duct is a low pressure system compared with portal channels.

In the first case, the patient had a very short history of portal hypertension and radiological findings clearly demonstrated by extrahepatic bile duct obstruction that resulted in intrahepatic biliary duct dilatation, sludge and calculi formation. Portal vein was totally thrombosed and CBD could not be found in its normal course as portal vein collaterals compressed it. Other two cases have prior history of chronic liver disease with secondary features of portal hypertension.

These are unusual presentations for cases of portal hypertension. Only few cases reported earlier had such grave presentation. For instance, Sumanthi et al presented a case of portal hypertension with obstructive jaundice secondary to portal biliopathy. Similarly, Chawla et al reported a case of portal biliopathy with obstructive jaundice and haemetemesis. The radiological findings in the current cases also suggested extensive periportal and peripancreatic varices, which caused almost complete compression of CBD. Moreover, intrahepatic dilatation with calculi formation was also seen in first case. So the radiological appearances of these cases are very typical of portal biliopathy as one does not often see the extrahepatic compression of CBD by collateral channels. This feature is not really seen in other causes of portal hypertension.

Proper case management is very important as prolonged biliary duct obstruction can lead to the development of ascending cholangitis or later on secondary biliary cirrhosis.

Management of portal biliopathy depends upon clinical manifestations. Asymptomatic patients generally do not require treatment. CBD calculi can be removed by endoscopic retrograde cholangiopancreatography (ERCP) and if there is stricture or narrowing in the duct, stent can be placed. Stones causing obstruction can be removed by endoscopic sphincterotomy.

Care should be taken, as there are large venous collaterals around ampullary region, which can bleed profusely. In patients with symptomatic biliary obstruction not amenable to endoscopic therapy, a Porto systemic shunt is indicated to decompress the portal system. If changes of portal biliopathy do not regress even then, hepaticojejunostomy can be performed to alleviate biliary obstruction.

Conclusion

Cavernous transformation of the portal vein due to extrahepatic portal vein obstruction is not infrequent but biliary obstruction in association with this disorder is distinctly uncommon. Proper case management is very important as prolonged biliary duct obstruction can lead to the development of ascending cholangitis or later on secondary biliary cirrhosis.

References