An Unusual Cause of Upper Gastrointestinal Bleeding: Duodenal Vascular Ectasia after Splenectomy

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ABSTRACT

Cavernomatous transformation of portal vein (CTPV) is characterized by a collateral network of veins in the hepatic hilum that bypasses an obstructed or occluded extrahepatic portal vein. We report a case of CTPV, 25 years after splenectomy in a 39-year-old man with acute and chronic blood loss from duodenal vascular ectasia without esophageal or gastric varices or portal hypertensive gastropathy. This case demonstrates that distal part of duodenum assessment is important in gastrointestinal bleeding of patients with CTPV.

Abbreviations: GI: Gastrointestinal; CTPV: Cavernomatous transformation of portal vein; CT: Computed tomography; PHD: Portal hypertensive duodenopathy.

Keywords: Cavernomatous transformation of portal vein, Gastrointestinal bleeding, Splenectomy, Vascular ectasia.


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INTRODUCTION

Duodenal vascular ectasia is a rare cause of upper gastrointestinal (GI) bleeding. It is associated with end-stage renal disease and aortic valve diseases, as well as a component of portal hypertension. Portal/splenic vein thrombosis, development of portal hypertension and portal vein cavernomatous transformation may be seen after splenectomy. Cavernomatous transformation of portal vein (CTPV) is new vascular network formation around gall bladder, intrahepatic and extrahepatic biliary tract as a result of portal vein obstruction. We presented a case with CTPV extending to duodenum while there were neither esophageal varices nor gastric varices.

CASE REPORT

A 39-year-old man with history of splenectomy and distal pancreatectomy because of a traffic accident 25 years ago was admitted to the hospital with the symptoms of fatigue, palpitation and intermittent melena for 3 months. He had a history of diabetes for 3 years, but no history of liver disease or alcohol abuse. On physical examination, he was pale; chest and heart examinations were normal. The abdomen was soft and there was no abdominal mass or bruit. Laboratory data showed that hemoglobin concentration was 8.6 gm/dl, erythrocyte count was 4.11 million/mm³, platelet count 434,000/mm³, alanine aminotransferase was 25 (20-55) U/l, aspartate aminotransferase was 24 U/l, gamma-glucuronyl transferase was 133 (53-150) IU/l, alkaline phosphatase was 139 (102-450) U/l, albumin was 4.7 (3.4-4.2) gm/dl, serum iron was 24 (32-283) µg/dl. On esophagogastroduodenoscopy, there were no esophageal or gastric varices but in the second part of duodenum vascular ectasia lesions was detected (Fig. 1). Endoscopic biopsy was performed on vascular ectatic mucosa and pathologic examination revealed focal lymphangiectasia and grossly dilates blood vessels between the muscularis mucosal and submucosal layers (Fig. 2). Local hemostatic agent was used to achieve hemostasis after biopsy and bleeding stopped without any complications. Dynamic computed tomography (CT) showed liver steatosis and CT angiography revealed CTPV and varicose vascular lesions around the duodenum connected with portal vein (Figs 3A to D). The patient was stabilized with supportive therapy. He was discharged 1 week after admission with propranolol prophylaxis. At 3-month follow-up, clinical assessment and laboratory data did not show any evidence of bleeding.

DISCUSSION

Noncirrhotic portal hypertension causes upper GI bleeding via esophageal or gastric varices and portal hypertensive gastropathy. This case demonstrates the importance of distal part of duodenum assessment in patients with CTPV.
Splenic vein thrombosis leads to a localized, left-sided or sinistral hypertension. Pancreatic surgery is another cause of splenic vein thrombosis. Isolated gastric varices result from thrombosis or obstruction of the splenic vein resulting in back pressure changes in the left portal system. The patient in this case report did not present with gastric varices or portal hypertensive gastropathy. CT angiography revealed only CTPV without splenic or portal vein thrombosis.

Cavernous transformation of the portal vein (also called portal cavernoma) in adults is an uncommon finding. In the pediatric age group, it is frequently associated to congenital anomalies. After thrombosis of the portal vein, in the absence of liver disease, organization and subsequent recanalization of the thrombus take place, followed by the development of hepatopedal portal collateral vessels, in order to restore portal perfusion, thus leading to the formation of a typical CTPV. The formation of these vessels is necessary in order to drain splenic and mesenteric venous flow into the intrahepatic portal branches and to maintain sufficient hepatic blood flow and normal hepatic function.
These hepatopedal collateral vessels look like cavernous angiomata when observed in gross specimen sections. CTPV usually presents with upper GI bleeding due to esophageal varices. CTPV may presents only with jaundice, due to compression of the common bile duct. In some cases, CTPV can favor the development of biliopathy. Whether the formation of portal thrombosis results in typical CTPV depends on the duration of portal obstruction. In our patient, detection of CTPV suggested the development of portal vein thrombosis after surgery. When portal obstruction is limited to the portal trunk only, the hepatopedal collateral veins of CTPV may skip the obstructed site and connect the intralobular open portal branch to maintain normal portal perfusion. In cases where portal obstruction is more extensive, even though the hepatopedal collateral veins of CTPV may take part in portal circulation, they are not sufficient to relieve portal hypertension, thus causing esophageal varices and formation of portacaval collateral circulation. Changes in intra-abdominal pressure during splenectomy decrease portal vein blood flow and induce stasis. Both stasis of venous flow and resulting congested coagulation factors may induce portal vein thrombosis. Stasis of blood in the stump of the splenic vein might be another mechanical factor for portal vein thrombosis. Turbulence in the stump of the splenic vein may result thrombosis and thrombus spread into portal vein. Although platelet count and hypercoagulability can increase postoperatively, association between these factors with formation of thrombosis at portal system is unclear. Thrombosis at portal system may be influenced by numerous technical factors, such as early ligation of the splenic artery, use of the endoscopic vascular stapler, or distal or proximal ligation of the splenic vein, but their definitive role is controversial. Interestingly, in this case, the extension of collateral veins to duodenum without varices suggested that portal obstruction was limited and collateral vessels at duodenum might be an anatomical variation or a consequence of technical factors at surgery. Only a few cases of vascular ectasia of the duodenum have been reported in patients with cirrhosis who had esophageal varices or portal hypertensive gastropathy. Duodenal lesions in portal hypertensive gastropathy patients are clinically significant because of being a possible source of massive bleeding. Barakat et al found portal hypertensive duodenopathy (PHD) in 51.4% of cirrhotic patients. Moreover, study have shown that PHD restricted to second part in duodenum was only 3.8% within PHD group and none of them had duodenal vascular ectasia. In the present case, there was bleeding from varicous collateral veins which originated from portal vein and extended to duodenum in the absence of portal hypertension findings at esophagus or stomach. To our knowledge, this is the first patient who has bleeding from isolated duodenal vascular ectasia with CTPV and noncirrhotic portal hypertension. Consequently, this case highlights the point that CTPV should be suspected in patients who had a history of splenectomy. Isolated duodenal vascular ectasia should be kept in mind as a rare cause of upper GI bleeding.

REFERENCES


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