Small Duct Sclerosing Cholangitis: A Case Report and Review of Literature

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Abstract: PSC often undiagnosed since approximately 40%-50% of patients with this disease are asymptomatic. Fatigue, fever, jaundice, pruritus, and vague upper abdominal discomfort are the most commonly described symptoms at the time of diagnosis. PSC involves both the intra- and extra-hepatic bile ducts in the majority of patients while an estimated 25% of patients only have isolated intra-hepatic involvement. In other reports intra- and extra-hepatic bile duct involvement occur in only 10% of cases.

Keywords: Sclerosing cholangitis, cholestasis, intra-hepatic, small ducts

Abbreviations: PSC: primary sclerosing cholangitis; SCC: secondary sclerosing cholangitis; NSAIDS: non steroidal anti-inflammatory drugs; HAV: hepatitis A virus; HBs: hepatitis B surface antigen; HBC: hepatitis B core; ANA: anti nuclear antibody; ASMA: anti smooth muscle antibody; AMA: anti mitochondrial antibody; ERCP: endoscopic retrograde cholangiopancreatography; CBD: common bile duct; AIH: autoimmune hepatitis; UC: ulcerative colitis.

1. INTRODUCTION

Primary sclerosing cholangitis (PSC) is a cholestatic liver and biliary tract disease caused by chronic inflammation of the biliary tract epithelium not attributed to any cause. This inflammation results in multifocal intra- and/or extra hepatic biliary strictures and fibrosis leading to biliary cirrhosis and malignancy development [1, 2]; we reported a case of cholestasis diagnosed as isolated intrahepatic sclerosing cholangitis or small duct PSC.

2. CASE PRESENTATION

Male patient 32 years old with no history of medical diseases apart from cholecystectomy 11 years ago began to complain from recurrent episodes of fatigue, upper abdominal pain, nausea, vomiting, pruritis and yellowish discoloration of skin associated with low grade fever, however, no diarrhea or bleeding per rectum is present. Patient sought medical advice and ultrasound examination was performed that revealed obstruction of common bile duct with missed stone, ERCP was performed and temporary stent was inserted for 3 month then removed, however patient remained to complain of jaundice and pruritis without improvement. No history of significant drug intake except for occasional NSAID for tooth pain. No family history of similar conditions. Examination was non significant apart from jaundice and epigastric tenderness with no lymphadenopathy, hepatosplenomegaly or neurological manifestations. Patient gave history of recurrent periods of bilirubin elevations followed by some drop of their levels in between. Initially patient's total bilirubin level was 14 and direct bilirubin was 7, alkaline phosphatase was 600 then dropped to 337 with drop of bilirubin level to 3.5 mg/dl, liver enzymes were normal, CBC revealed leucocytosis of about 14,000 with normal levels of hemoglobin, platelets, esinophil counts as well as normal bleeding profile. Viral markers as HAV IGM, HAV IGg, HBs antigen, HBC IGM, HBC IGg, HCV antibody were all negative. ESR 1st hour 38 and 2nd hour 71 , autoimmune markers as ANA, ASMA, and AMA were negative. Pelvi abdominal ultrasound shows average size liver and spleen, no ascitis and normal caliber of CBD and intrahepatic biliary radicals that was confirmed with triphasic abdominal CT that was also free apart from insicional postcholecystectomy hernia. Liver biopsy was asked and revealed that liver tissue shows bile stasis with dense bile pigment in distended hepatocytes. Sinusoids are patent. Portal tract shows dense sclerotic bands,
inflammatory cells, cholangitis and fibrous tissue amidst bile ducts. No evidence of regeneration nodules or cirrhotic changes, viral inclusions inside hepatocytes, dysplastic or malignant changes or features of autoimmune hepatitis (AIH) or overlap syndrome.

Figure 1(A): Classic ‘onion skin’ fibrosis, causing obliteration of the bile ducts in primary sclerosing cholangitis [Photograph courtesy: Thomas Smyrk, M.D., Department of Pathology, Mayo Clinic, Rochester]

Figure 1(B): Ductopenia, without onion-skinning fibrosis in small-duct primary sclerosing cholangitis [Photograph courtesy: Thomas Smyrk, M.D., Department of Pathology, Mayo Clinic, Rochester]

3. DISCUSSION

Patients with biochemical markers and histological features suggestive of PSC with normal cholangiography findings are considered to have small-duct PSC (3). Small duct PSC may be an early stage of PSC (4). Small-duct PSC is a condition which affects only microscopic bile ducts (i.e., bile ducts seen only on liver biopsy) (5). Many patients with PSC are asymptomatic with no physical abnormalities at presentation. The diagnosis is made incidentally when persistently cholestatic liver function tests are found (6). This subgroup of PSC had been less studied than the classic patients with large-duct PSC. Gallbladder wall thickening or enlargement (7) gallstones, cholecystitis, and mass lesions were identified in up to 41% of patients with PSC who undergo ultrasound examinations (8). Serum aminotransferase levels are elevated in the majority of patients with this disease (about 2–3 times upper limits of normal), but like the alkaline phosphatase can also be in the normal range (9). CT imaging can detect thickening of the bile ducts with contrast enhancement consistent with inflammation, saccular dilatations of the intrahepatic ducts, heterogeneous bile duct dilatation, document the presence of portal hypertensive complications (i.e., varices, splenomegaly, and ascites), and can identify mass lesions (10-14). Detectable auto antibodies are found in about 97% of patients with PSC (15), these auto antibodies, however, are not routinely used for the diagnosis of PSC, as they may not be present and furthermore do not correlate with disease severity or disease prognosis (16). The classic histopathologic finding of PSC is presence of periductal concentric (“onion-skin”) fibrosis, but this observation is infrequent in liver biopsy specimens from patients with PSC and may also be observed in SSC (17). Better prognosis in individuals with small-duct PSC was documented in many studies, however, data on this subgroup of PSC is limited due to lack of long-term follow up (18,19). A recent study for patients with small duct PSC suggested that approximately 25% of patients progress to classic PSC in an average of 8 years during their follow up (20). Individuals with small-duct PSC may progress to end-stage liver disease even without developing large duct disease. Cholangiocarcinoma however, does not seem to occur in patients with small-duct disease without progression to large-duct PSC (20). The association of small-duct PSC appears to be stronger with Crohn’s colitis than with UC (21, 22), however as put by Jurgen Ludwig [23], the distinguished liver pathologist formerly at the Mayo Clinic: “No valid scientific reason can be found to suggest that small-duct PSC could not occur without large-duct PSC and IBD. Ursodeoxycholic acid (UDCA) is widely used in the treatment of a variety of cholestatic liver diseases, but very limited data exist on its effects in small-duct PSC (24). No controlled trials of medical therapy including only patients with small-duct PSC have been performed. Patients with large-duct PSC were shown to have a better response to immunosuppressive therapy than patients with small-duct PSC (25). The better results of immunosuppressive therapy in the large-duct patients were probably linked to higher inflammatory activity indicated by the higher ALT levels in the large-duct patients as compared with the levels in the small-duct group (25). For the most part, the course of disease is milder than for large-duct disease with fewer symptoms and a favorable prognosis with regard to survival, need for transplantation is
extremely small (26). Small-duct PSC may recur after liver transplantation (5)

Figure 2: Microscopic Picture for Liver Biopsy Obtained from our Patient

4. PATIENT CONSENT FORM

Informed consent was taken from concerned patient.

REFERENCES


