Presentation of Intrahepatic Cholangiocarcinoma with Fulminant Hepatic Failure: A Case Report

Thara Pratap1  Muhammed Jasim Abdul Jalal2  Pushpa Mahadevan3  Abraham Koshy4  Roy J. Mukkada4  Pradeep G. Mathew4  Rithu Sebastian4

1Department of Radiology, VPS Lakeshore Hospital, Kochi, Kerala, India
2Department of Internal Medicine and Rheumatology, VPS Lakeshore Hospital, Kochi, Kerala, India
3Department of Pathology, VPS Lakeshore Hospital, Kochi, Kerala, India
4Department of Gastroenterology, VPS Lakeshore Hospital, Kochi, Kerala, India

Fulminant hepatic failure as initial presentation due to diffuse parenchymal infiltration by cholangiocarcinoma is a rare entity. We present the case of a 49-year-old female patient who had a fatal outcome with acute liver failure due to diffuse intrahepatic cholangiocarcinoma. No definite mass lesion was identified on cross-sectional imaging. The final diagnosis was made on transjugular liver biopsy. This discussion highlights the possibility of infiltrative cholangiocarcinoma as a rare cause of fulminant hepatic failure.

Abstract

Keywords
► fulminant hepatic failure
► intrahepatic cholangiocarcinoma
► transjugular liver biopsy

Introduction

Intrahepatic cholangiocarcinoma (ICC) is the second most common malignant tumor of the liver. It constitutes 10% of cholangiocarcinomas.1 By definition, intrahepatic cholangiocarcinoma originates proximal to second degree bile ducts. This tumor is classified on the basis of gross morphological features into mass forming, periductal infiltrating, and intraductal type. Cholangiocarcinoma often presents at an advanced stage and most of the cases are unresectable. The disease often has a fatal outcome. The prevalence of this disease is highest in Southeast Asian countries such as Thailand and is less in the Western world. In Asian countries, parasites such as liver fluke—Clonorchis sinensis and Opisthorchis viverrini—and hepatolithiasis are major risk factors, other causes being biliary tract disease such as primary sclerosing cholangitis, primary biliary cirrhosis, choledochal cysts, infective causes such as hepatitis B and C virus, alcoholic liver disease, and smoking, though most of the cases are idiopathic. The disease is more common in males than females and occurs most frequently between sixth and seventh decade. The incidence of the disease is on the rise in the west2 and in younger population.

Case Presentation

A 49-year-old lady presented with recurrent dull aching pain in the right upper abdomen and jaundice of 1 month duration. It was associated with dyspnea on exertion. On examination, she was icteric and had abdominal distension with shifting dullness, bilateral pedal edema, and facial puffiness. There was no history of pruritus, clay colored stools, weight loss, fever, or prodromal symptoms. She was on ayurvedic medications for urinary tract infection 3 months back. Her liver function (AST/ALT/ALP 92/43/331) was deranged and coagulation parameters (PT/INR/Index 15.2/1.13/90.8%) were found to be normal. Viral markers were nonreactive. Ascitic fluid analysis was lymphocyte predominant with negative cytology. Ascitic fluid albumin was normal. Upper gastrointestinal endoscopy showed grade II esophageal varices and small fundic varix suggesting a diagnosis of decompensated liver disease.
Cross Sectional Imaging

Triple phase computed tomography (CT) scan was performed with GE 64 slice scanner light speed VCT XTE. Ninety milliliter of intravenous contrast Ultravist 350 mg was given at a flow rate of 3.8 mL/min. CT images were obtained with bolus tracking technique with a scan delay of 4 to 5 seconds after a set contrast threshold of 100 HU. Arterial, porto-venous, hepatic venous and delayed scans were taken. The images were viewed on a dedicated GE workstation (AW 4.4) and coronal, sagittal reconstructions were also analyzed.

Contrast-enhanced computed tomography (CECT) scans showed hepatomegaly with patchy ill-defined hypo-dense areas involving both lobes better appreciated in portal and venous phase scans (►Figs. 1–4) than in arterial phase. No definite enhancing mass lesions, ductal thickening, or biliary

Fig. 1 (A–F, arterial phase): 64 slice contrast CT scans showing hepatomegaly with lobulated contour of liver with patchy hypo-dense areas involving both lobes of liver. CT, computed tomography.

Fig. 2 (A–F, portal venous phase): contrast CT scans showing hepatomegaly with multiple patchy geographic hypo-dense areas involving both lobes of liver, better appreciated than in arterial phase scans. CT, computed tomography.
dilatation was noted. There was associated small segment portal vein thrombosis involving right posterior segmental branch. Rest of the portal vein branches was normal. In addition, there were minimal ascites and multiple small retroperitoneal nodes. Differential diagnosis considered was chronic liver disease, infiltrative liver disease, and metastatic liver disease. However, there was no evidence of primary in the chest or abdomen.

In view of acute liver failure of unknown etiology, transjugular liver biopsy was performed. Histopathology revealed the diagnosis of infiltrating periductal adenocarcinoma with immunohistochemistry confirming the diagnosis of primary cholangiocarcinoma liver. Oncology evaluation was sought, but she deteriorated rapidly.

**Histopathology**

The tumor was of periductal type with multiple foci of infiltrating adenocarcinoma predominantly of clear cell type (Figs. 5 and 6). Clear cell changes have been uncommonly reported in patients with cholangiocarcinoma.3

---

**Fig. 3** (A–F, hepatic venous phase): contrast CT scans show hepatomegaly with patchy hypo-dense areas involving both lobes. (D) Shows small segment portal venous thrombosis. CT, computed tomography.

**Fig. 4** (A–F, delayed scans): further iso-dense appearance of lesions is shown.
Intrahepatic Cholangiocarcinoma with Fulminant Hepatic Failure

Pratap et al.

Immunohistochemistry showed tumor cells positive for cytokeratin 7 (CK7) and negative for hepatocyte antibody (Figs. 7 and 8).

Discussion

ICC with typical imaging findings can be diagnosed easily. Our patient presented with hepatomegaly with patchy ill-defined hypo-dense areas and no definite mass on cross sectional imaging. In spite of periductal infiltrating type on histopathology, there was no biliary duct thickening or bile duct obstruction. This is an extremely rare presentation on imaging.

Pathology showed extensive periductal type of disease explaining the cause of acute liver failure, which is often due to neoplastic infiltration of hepatic sinusoids leading to parenchymal infarction and secondary necrosis of hepatocytes. Replacement of 80 to 90% of hepatic parenchyma by neoplastic cells could lead to jaundice and liver failure.

Literature search reveals only few cases of primary cholangiocarcinoma with acute liver failure. The diagnosis in these cases was made on liver biopsy or autopsy. In none of the cases, preoperative diagnosis could not be made as no definite masses could be identified on CT scan.

Fulminant hepatic failure due to primary liver tumors other than hepatocellular carcinoma is very rare. There are few case reports of acute liver failure due to primary hepatic...
Intrahepatic Cholangiocarcinoma with Fulminant Hepatic Failure
Pratap et al.
Journal of Gastrointestinal and Abdominal Radiology ISGAR Vol. 2 No. 2/2019

angiosarcoma. Infiltrative deposits due to hematopoietic diseases and metastatic deposits from primary breast and lung cancers are already known entities.

Conclusion
Cholangiocarcinoma presenting as diffuse infiltrating intrahepatic disease with no definite mass lesions on CT is very rare. The role of liver biopsy is crucial to make the diagnosis given the atypical imaging findings and presentation. This alerts us to the rare possibility of primary diffuse infiltrating cholangiocarcinoma as a cause of fulminant hepatic failure.

Conflict of Interest
None declared.

References