A rare cause of biliary obstruction – intraductal neuroendocrine tumor of the right hepatic biliary duct: a case report

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Abstract

Primary biliary tract neuroendocrine tumors (NETs) are extremely rare tumors that account for 0.2-2% of all gastrointestinal neuroendocrine tumors. The typical presentation is with jaundice and other symptoms related to biliary obstruction. We present a case of right hepatic duct NET in a 27-year-old female patient, asymptomatic, presented for a routine ultrasound examination that revealed moderate dilatation of the intrahepatic biliary ducts and a 20 mm hyperechoic lesion in the right hepatic biliary duct. Additional imaging was performed with the presumptive diagnosis of cholangiocarcinoma. After surgery, the histopathological and immunohistochemical report was conclusive for the diagnosis of G2 well-differentiated NET.

Keywords: neuroendocrine tumor; right hepatic bile duct; biliary obstruction

Introduction

Primary biliary tract neuroendocrine tumors (NETs) are extremely rare tumors with less than 100 reported cases in the literature [1,2], accounting for 0.2-2% of all gastrointestinal neuroendocrine tumors [2,3]. NET is derived from embryonal neural cells called Argentaffin or Kulchitsky-enterochromaffin cells that have the potential of secreting serotonin. These cells are found in the highest proportion in the small intestine and rarely within the biliary ducts, explaining the low incidence of extrahepatic biliary neuroendocrine tumors (NET) [2,4-6].

According to the World Health Organization (WHO) classification system (2010) NETs are defined as neoplasms with neuroendocrine differentiation. WHO classifies neuroendocrine neoplasms based on their Ki67 and Mitotic indices. In 2017, the WHO classification was updated based on recent evidence [7-9].

We report a case of primary biliary NET arising from the right hepatic biliary duct. After an extensive search of the literature, no references regarding a NET of the right hepatic biliary duct was found.

Case report

A 27-year-old female patient, asymptomatic, with no known pathology, presented for the evaluation of a minor cytolytic syndrome. Clinical examination revealed no hepatomegaly/splenomegaly, jaundice or other abnormalities. Laboratory tests – slightly increased transaminases: AST 40 U/L (N<35), ALT 64 U/L (N<35) with negative markers of viral B or C chronic hepatitis. Diagnostic workup with an abdominal ultrasound revealed moderate dilatation of the intrahepatic biliary ducts and a 20 mm hyperechoic lesion in the right hepatic biliary duct (fig 1). Contrast enhanced ultrasound (CEUS) examination showed a homogenous hyperenhancement pattern in the arterial phase with intense washout in the late phase, suggesting malignancy (fig 2).
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Additional laboratory tests showed: mild cholestasis AP 206 U/L (N<129), γ-glutamyl transpeptidase 286 U/L (N<38). Tumor markers, including carcinogenic embryonic antigen (CEA), carbohydrate antigen 19-9 (CA19-9) and alpha-fetoprotein, were normal.

Subsequently, abdominal MRI and MRCP confirmed a solid lesion in the right hepatic biliary duct, with discreet hyperintensity in T2 and restricted diffusion. In the delayed phase, the lesion presented washout, suggestive of malignancy (fig 3).

The presumptive diagnosis was biliary obstruction due to localized Klatskin/cholangiocarcinoma tumor without evidence of metastasis.

The patient was referred to the Department of Surgical Oncology and resection of the common bile duct, cholecystectomy, Roux-en-y hepatico-jejunostomy reconstruction and locoregional lymphadenectomy was performed. The histopathological and immunohistochemical report of the resected tumor was conclusive for the diagnosis of a well-differentiated neuroendocrine tumor (NET G2). Immunohistochemical staining showed that the tumor cells were strongly positive for Chromogranin A and Synaptophysin with the Ki-67 index 3-5%.

After surgery, the patient received chemotherapy and has remained under oncologic surveillance. There was no recurrence of the disease at three years after surgery.

Discussion

Neuroendocrine tumors derive from Argentaffin or Kulchitsky-enterochromaffin cells. The most common sites of NET occurrence are ileum, appendix, bladder, prostate, rectum, stomach, bronchi, pancreas, and biliary tree. The paucity of enterochromaffin cells in the biliary tree explains the rare incidence of NETs in the biliary system. Biliary tract NETs are extremely rare and only one-fifth of these tumors are well-differentiated NETs [5,10]. Biliary tract NETs are often non-secreting tumors and larger tumors over 2 cm are associated with aggressive behavior [5].

Published data show that in most cases the presenting symptom was related to the local invasion of the tumor [10]. The most common presenting symptom in patients with biliary tract NETs is jaundice, followed by pain and other nonspecific symptoms (pruritus, nausea, vomiting, weight loss) [1,11]. Our patient was asymptomatic probably due to the small dimensions of the tumor.

Cholangiocarcinoma is the most frequent biliary tract malignancy. Differential diagnosis between cholangiocarcinoma and other bile duct tumors, such as NET, is challenging before surgical resection and histopathological exam. In our case, the final diagnosis was also established postoperatively.

NETs of the biliary ducts are generally well-differentiated, slow-growing tumors with rare metastatic spread and have a favorable prognosis with a high-rate survival, mostly in cases in which curative surgical resection is possible [11-15]. Our patient also had a well-differentiated biliary tract NET- G2.

Fig 1. Abdominal ultrasound showing moderate dilatation of the intrahepatic biliary ducts and a slightly hyperechoic lesion, 20 mm in size, in the right hepatic biliary duct.

Fig 2. CEUS examination: a) homogenous hyperenhancement pattern in the arterial phase; b) mild washout starting in the portal vascular phase; c) obvious washout in the late vascular phase.
The literature search did not reveal other cases of NET isolated in the right hepatic biliary duct, possibly making our case the first reported case of an intraductal NET of the right hepatic biliary duct.

In conclusion, biliary tract NETs have as a typical presentation, jaundice and other symptoms related to biliary obstruction, but patients can also be asymptomatic. Due to the absence of specific symptoms, a correct preoperative diagnosis is rare: diagnosis is usually made postoperatively, based on the histopathology exam. Radical tumor surgery is the only available curative approach, with high survival rates.

References