IgG4 Associated Cholangitis and Hepatic Pseudoinflammatory Tumour: IgG4 Related Disease Mistaken by Neoplastic Disease

Colangite e Pseudotumour Inflamatório Hepático Associados a IgG4: Doença Associada a IgG4 Simulando Doença Neoplásica

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Abstract
We report the case of a 71-year-old man presenting inaugural cholestatic jaundice, with a stricture and circumferential wall thickening of the distal common bile duct, as well as an ill-defined hepatic infiltrative mass with “amputation” of peripheral distended bile ducts. The patient underwent a pancreaticoduodenectomy and hemihepatectomy for suspected cholangiocarcinoma. Histology revealed IgG4 associated cholangitis (IgG4-AC) and hepatic pseudoinflammatory tumour (IPT). IgG4-related disease (IgG4-RD) is a recently established systemic disease, characterized by fibro-inflammatory tissue infiltrates and tumefaction of one or more organs. The hepatobiliary manifestations comprise IgG4-AC and associated IPT - which can be mistaken for cholangiocarcinoma. Clinicians must be familiar with this disorder, usually responsive to corticosteroid therapy, in order to avoid unnecessary invasive procedures. We present a case of IgG4-AC and associated IPT and review the literature of this rare presentation of IgG4-RD.

Keywords
IgG4-related disease; Autoimmune cholangitis; Hepatic pseudoinflammatory tumour; Cholangiocarcinoma; Hemihepatectomy; Cephalic pancreaticoduodenectomy.

Introduction
Immunoglobulin G4-related disease (IgG4-RD) is a systemic condition affecting virtually every organ system that comprises common pathologic, serologic and clinical features. It has been increasingly recognized since 2003, when Kamisawa et al. proposed that a group of disorders, previously thought to be unrelated, might be part of a broader spectrum of IgG4-RD; nevertheless, pathogenesis is still poorly understood. Common features include tissue infiltration by IgG4-positive plasma cells and lymphocytes, as well as a variable degree of fibrosis and tumour-like swelling in one or more organs. Symptoms are mild, with neither fever nor elevation of C-reactive protein levels. Diagnosis requires meticulous correlation of the histopathologic features with clinical and radiologic findings. Imaging plays an important role in demonstrating enlargement of involved organs, detecting multiorgan involvement and, eventually, for assessing response to therapy.

Clinical Case
We report the case of a 71-year-old man who presented himself at the emergency department of our hospital in August 2015 complaining of abdominal pain for the previous week, jaundice, choluria and diarrhea with acholic faeces. At physical examination, the abdomen was tender at the epigastrium and right upper abdomen. The patient had a previous clinical history of bladder neoplastic polyps as low grade papillary urethral non-invasive carcinoma removed by transurethral resection in 2008 and 2010, diabetes mellitus type 2, arterial hypertension and vascular peripheral disease and he was regularly medicated with metformin and...
vildagliptin, enalapril and hydrochlorothiazide, diosmin and esomeprazole.

Blood tests at admission showed high levels of total bilirubin 9.90mg/dL (<1.2mg/dL) with a predominant increase of the direct bilirubin (conjugated component), as well as high levels of alanine aminotransferase 116mg/dL (12-78mg/dL), aspartate aminotransferase 122mg/dL (<34mg/dL), gamma-glutamyl transferase 1330mg/dL (<34mg/dL), alkaline phosphatase 880mg/dL (40-130mg/dL); slightly elevated C-reactive protein 0.9mg/dL (<0.5mg/dL); abnormal presence of moderate quantity of bilirubin in urine analysis; low levels of amylase (3 U/L) and haemoglobin of 11.8 mg/dL (13-17.5mg/dL). Further workup, during the hospital stay, revealed normal levels of the tumour markers CEA of 1.9mg/dL (<5mg/dL) and CA 19.9 of 23.4mg/dL (0-37mg/dL), as well as further progressive increase of total bilirubin level.

An abdominal ultrasound was performed at admission (Fig. 1), which revealed bilateral dilatation of the intrahepatic biliary ducts, as well as the proximal half of the common bile duct (CBD), with stenosis and a significant hypoechoic circumferential wall thickening of the remaining CBD; non distended gallbladder, without signs of cholelithiasis; diffuse atrophy of the pancreatic parenchyma and mild dilatation of the main pancreatic duct.

A computed tomography (CT) was performed subsequently (Fig. 2), confirming the previous findings of biliary ducts dilatation (CBD calibre of 19mm), with stenosis and a significant circumferential and regular wall thickening of the distal common bile duct, with homogeneous contrast enhancement; also, diffuse atrophy of the pancreatic parenchyma, with some calcific foci, mild diffuse dilatation (4-5mm) and slight irregularity of the main pancreatic duct (suggestive of chronic pancreatitis). These findings were

Figure 1 – Ultrasound. (A) Axial plane, showing bilateral dilatation of the intra-hepatic biliary ducts (arrows). (B) Axial plane and (C) sagittal plane, demonstrating dilatation of the proximal CBD (arrow), with stenosis and hypoechoic circumferential wall thickening of the remaining CBD (open arrows). (D) Axial plane, showing diffuse atrophy of the pancreatic parenchyma and mild dilatation of the main pancreatic duct (arrows). It was also noticed moderate hepatomegaly with signs of steatosis and minor splenomegaly.

Figure 2 – Enhanced computed tomography (CT). (A) Axial plane in a liver dedicated window, depicting a poorly circumscribed area of heterogeneity on segment II of the left liver lobe, with mild delayed enhancement (open arrows), causing “amputation” of peripheral dilated intra-hepatic bile ducts (arrows). Axial plane at a higher level (B) and a lower level (C), as well as coronal plane (D), demonstrating dilatation of the bile ducts (arrows), with stenosis and a significant circumferential and regular wall thickening of the distal CBD, with homogenous contrast enhancement (open arrows). Also, notice the diffuse atrophy of the pancreatic parenchyma, with some calcific foci, and mild diffuse dilatation and slight irregularity of the main pancreatic duct.
The magnetic resonance cholangiopancreatography, also confirmed the previous findings and further revealed an area of left hepatic parenchyma heterogeneity with ill-defined margins at segment II, presenting mild hyperintensity at T2-weighted images (WI) and hypointensity at T1-WI, associated with "amputation" of peripheral dilated intrahepatic bile ducts and an altered luminal signal intensity (hypointensity at T2-WI and hyperintensity at T1-WI) in the proximal dilated bile ducts. Retrospectively, a corresponding ill-defined infiltrative mass, with a mild delayed enhancement, was also depicted on CT.

Based on the clinical and imaging elements, a presumptive diagnosis of multifocal cholangiocarcinoma was made. Since no alternative diagnosis was proposed, biopsy was not performed and the patient underwent surgery. A cephalic pancreaticoduodenectomy and a left hemihepatectomy were performed. The pathological findings were highly suggestive of IgG4-related disease, with IgG4-AC and a hepatic pseudoinflammatory tumour (with approximately 4cm), probably related to a focal IgG4-AC exacerbation; no neoplastic tissue was found. Ensuing blood tests revealed IgG values of 2160 mg/dL, but the IgG4 serum levels were not subsequently assessed.

The post-operative course was complicated by a surgical wound infection, a fluid collection adjacent to the jejunopancreatic anastomosis, another infected haematic collection in the left upper abdomen, as well as a left empyema, that required intravenous broad spectrum antibiotics and percutaneous drainage. Except for the surgical wound infection, these complications were, eventually, managed after a long course of conservative treatment and the patient was discharged home 39 days after surgery. The surgical wound infection went on for several months, postponing the beginning of treatment with systemic corticosteroids.

Discussion

The hepatobiliary manifestations of IgG4-related disease (IgG4-RD) comprise cholangitis, pseudoinflammatory tumour and IgG4 hepatopathy. After the pancreas, the bile ducts are the second most commonly involved organ, seen in up to 88% of patients with IgG4-RD. IgG4-associated cholangitis (IgG4-AC) occurs more frequently in patients with type 1 autoimmune pancreatitis (60-70%), but it might also occur without pancreatic involvement, making the diagnosis more
difficult.\textsuperscript{5,6} Involvement by other extrapancreatic lesions may be additionally seen.\textsuperscript{2,6} Generally, IgG4-AC affects older patients and can induce obstructive jaundice.\textsuperscript{3} Both intra- and extrahepatic segments can be affected by dense bile duct infiltration of IgG4-positive plasma cells and extensive fibrosis, but the most commonly involved segment is the intrapancreatic portion of the common bile duct (CBD). The typical imaging appearance, at MRCP, is focal or diffuse bile duct wall thickening - usually long and continuous strictures are depicted - mostly associated with stenosis and upstream dilatation.\textsuperscript{7} Isolated strictures of the distal CBD may also occur.\textsuperscript{4} CT and MR imaging usually depict a circular and symmetric ring of tissue encasing the bile duct wall, with relatively smooth margins and homogeneous enhancement in the delayed phase.

The main differential diagnoses are cholangiocarcinoma and primary sclerosing cholangitis (PSC). The latter would not be expected in this case due to the acute onset of symptoms, older age and no previous history of inflammatory bowel disease; also, the typical findings for PSC are multifocal and short intrahepatic biliary strictures with beaded or “pruned-tree” appearance.\textsuperscript{8} Differentiation from cholangiocarcinoma may be difficult, especially in the presence of a soft-tissue mass that produces stenosis of the hilar hepatic bile duct, presented as inaugural jaundice. In the case of cholangiocarcinoma, tumour infiltration is usually confined to either extrahepatic or intrahepatic bile ducts, whereas in IgG4-AC luminal irregularities and stenosis generally involve both the biliary and pancreatic ducts.\textsuperscript{9} Differentiating hilar cholangiocarcinoma from other causes of hilar obstruction remains an issue.\textsuperscript{6} There are a few reports in the literature addressing cases of IgG4-cholangitis of the CBD, with no apparent pancreatic involvement, that underwent surgery for suspected cholangiocarcinoma.\textsuperscript{10-12}

IgG4 related cholangitis and hepatic inflammatory pseudotumour (HIP) is an extremely rare association, in which the cholangitis is considered either as the condition preceding or associated with the inflammatory pseudotumour.\textsuperscript{13} Inflammatory pseudotumours occur most commonly in the lung and only occasionally in extrapulmonary organs, including the liver in about 8% of cases.\textsuperscript{4} Inflammatory pseudotumours are characterised histologically by the proliferation of fibroblasts or myofibroblasts and inflammatory cell infiltrate, independently of the organs where they occur. Infections and autoimmune reactions have been suggested as possible triggers, although pathogenesis of hepatic inflammatory pseudotumours is still incompletely understood and might differ among cases.\textsuperscript{10} On CT, homogenous delayed enhancement was considered an important diagnostic imaging characteristic of IgG4-related disease. On MRI, these lesions are typically hypointense on T1-WI and hyperintense on T2-WI, but contrast enhancement patterns are variable.\textsuperscript{13} The differential diagnosis is generally with a primary liver neoplasm or metastasis, if the bile ducts or pancreas are also involved by the inflammatory disease. The IgG4-RD usually demonstrates a favourable response to corticosteroid therapy. However, it is strongly recommended to confirm the diagnosis of IgG4-RD by a biopsy specimen instead of a steroid trial, since some patients are refractory to such therapy and some malignant lesions may improve after steroid treatment. Relapses may sometimes occur, even under maintenance of corticosteroid therapy;\textsuperscript{12} azathioprine is frequently used in these cases.\textsuperscript{10} In the case we present, there was a lymphoplasmacytic infiltrate enriched in IgG4-positive plasma cells at histology, involving both the intra- and extrahepatic bile ducts. However, at imaging, no luminal irregularities or stenosis of the intrahepatic bile ducts were seen, but only a dominant stricture of the distal CBD causing upstream dilatation. This showed typical circumferential regular thickening of the wall and homogeneous enhancement. There were also pancreatic findings of chronic pancreatitis (confirmed at histology). Additionally, a probable exacerbation of the cholangitis at the left hepatic lobe caused a “mass-like” appearance, related to an inflammatory pseudotumour. The finding of a hepatic “mass” coupled with CBD stenosis and thickening with upstream dilatation may mimic multifocal cholangiocarcinoma, which represents about 5% of cholangiocarcinomas.\textsuperscript{16} The associated presentation of IgG4-AC and HPT is extremely rare and was misleading. The older age of the patient and the inaugural presentation with obstructive jaundice did not help in establishing the differential diagnosis. Furthermore, tumour markers CA 19.9 and CEA are not elevated in 15% and 70% of cholangiocarcinomas, respectively.\textsuperscript{16}

In conclusion, differentiating IgG4-related hepatobiliary disease from cholangiocarcinoma remains a challenge. Altogether, we believe that this was an extremely difficult case to differentiate from cholangiocarcinoma, based on clinical presentation and imaging alone. Therefore, we expect to emphasize the need for a biopsy specimen before potentially harmful and unnecessary invasive procedures are undertaken. Moreover, we suggest regular testing on IgG4 blood levels when there are overlapping hepatobiliary imaging findings, bearing in mind that it is neither specific nor uniformly increased in all patients, but it might guide clinicians into the right track for IgG4-RD diagnosis.

References