CLINICAL CASE

Autoimmune pancreatitis and ulcerative colitis: A clinical challenge of a true association

Pedro Barreiro¹, Pedro Pinto Marques², Gilberto Couto², David Serra¹,², Cristina Chagas¹, Leopoldo Matos¹

¹ Department of Gastroenterology, Egas Moniz Hospital, Lisbon, Portugal
² Digestive Disease Center, Luz Hospital, Lisbon, Portugal

Received 21 July 2011; accepted 5 October 2011
Available online 31 July 2012

KEYWORDS
Autoimmune pancreatitis; Chronic pancreatitis; Ulcerative colitis; Inflammatory bowel disease; Immunosuppression

Abstract Autoimmune pancreatitis is emerging as a well-defined clinical entity, yet its diagnosis and therapeutic approach still constitute a clinical challenge. Its association to other autoimmune diseases, namely ulcerative colitis, is known although the exact relationship between the two entities is not completely clarified.

We present the case of a patient who developed obstructive jaundice with later onset of blood-stained diarrhea, leading to a final diagnosis of autoimmune pancreatitis and ulcerative colitis.

We make a brief revision of autoimmune pancreatitis, its relationship with ulcerative colitis and therapeutic approaches of the same, namely the eventual necessity of immunosuppressive therapy.

© 2011 Sociedade Portuguesa de Gastrenterologia Published by Elsevier España, S.L. All rights reserved.

PALAVRAS-CHAVE
Pancreatite auto-imune; Pancreatite crónica; Colite ulcerosa; Doença inflamatória intestinal; Imunosupressão

Pancreatite auto-imune e colite ulcerosa: um desafio clinico de uma associação real

Resumo A pancreatite auto-imune é compreendida cada vez mais como uma entidade clínica bem definida, contudo o seu diagnóstico e abordagem terapêutica constitui ainda um desafio clínico. A sua associação com outras doenças auto-imunes, nomeadamente com a colite ulcerosa é conhecida porém a verdadeira relação entre as duas entidades não está totalmente esclarecida.

Apresentamos o caso clínico de um doente que iniciou quadro de icterícia obstrutiva chegando-se ao diagnóstico final de pancreatite auto-imune. Durante a investigação clínica o doente apresenta quadro de diarreia sanguinolenta diagnosticando-se colite ulcerosa extensa associada.
Introduction

For decades, different cases of chronic pancreatitis associated with an important immunological component have been recognized. In 1961, Sarles et al. used the term "primary inflammatory pancreatitis" to describe a group of patients with pancreatitis, until then of unknown aetiology, who presented little or no abdominal pain, cholestasis, increased serum immunoglobulins and severe pancreatic inflammatory infiltrate with fibrosis. Since then many terms were employed to describe cases of pancreatitis with similar characteristics until 1995 when, for the first time, the term autoimmune pancreatitis (AIP) was applied. From this date, many advances in the understanding of this entity have been recorded.

At the same time, an increased incidence of pancreatic diseases in patients with inflammatory bowel disease (IBD) has been reported, namely with ulcerative colitis (UC). This may be drug-related or due to the increased incidence of cholelithiasis among IBD patients. However rarer forms of chronic pancreatitis are described, and its association with AIP is underlined by different case reports, although the true incidence is still unknown.

Clinical case

We present the case of a 34-year-old white man with no past medical history who developed malaise, fatigue, persistent epigastric discomfort and one month later jaundice. There was no history of alcohol intake, drug abuse or medication. The physical exam was unremarkable except for jaundice and epigastric pain. Laboratory evaluation was remarkable for abnormal liver function tests with cholestasis and slight hepatic cytolysis (alkaline phosphatase, 340 UI/L; gamma-glutamyl transferase, 191 UI/L; total bilirubin, 5.57 mg/dl; aspartate aminotransferase, 86 UI/L; alanine aminotransferase, 102 UI/L). Abdominal ultrasound was consistent with extra-hepatic cholestasis and an abdominal computed tomography (CT) documented common bile duct (CBD) narrowing at the pancreatic level, which was described as normal. The endoscopic retrograde cholangiopancreatography (ERCP) confirmed the intra-pancreatic regular CBD stenosis without further changes of the extra-pancreatic bile structures (Fig. 1A). Biliary citology was negative for malignancy. Pancreatic duct cannulation was unsuccessful and a 10 Fr biliary stent was placed (Fig. 1B).

For further evaluation a magnetic resonance imaging-cholangiopancreatography (MRI-CP) was ordered, which revealed discrete pancreatic head heterogeneity, with no
main pancreatic duct (MPD) abnormalities. An endoscopic ultrasound (EUS) showed an abnormal pancreatic head, overall hypoechoic, heterogeneity and slightly increased, with no MPD visualization (Fig. 2). This was felt suggestive of AIP and fine needle aspiration with a 19 g Trucut needle (Cook) at the pancreatic neck was performed. Histology showed extensive pancreatic fibrosis, marked ductopenia, diffuse lymphocytic infiltration predominantly periductal as well as peri-venular lymphocytic infiltrates (Fig. 3). These findings were felt to support the diagnosis of AIP.

Additional laboratory evaluation showed increase of IgG4 (212 mg/dl). The autoantibodies studied (ANA, AMA, Ac Anti-DNA, ASMA, ANCA e ASCA) and the rheumatoid factor were normal.

One week after the initial diagnostic workup, the patient presented with blood-stained diarrhea and abdominal pain. Stool culture, stool evaluation for ova and parasites and Clostridium difficile toxin assay were negative. Colonoscopy showed diffuse continuous superficial erosions and ulcerations throughout the entire colon and rectum with loss of the vascular pattern. Histology supported the hypothesis of active UC diagnosis.

The clinical, analytical, imaging and histological evaluation of the patient therefore allowed for establishing the diagnosis of AIP associated with UC.

The patient was started on prednisolone 40 mg qd for 2 weeks combined with mesalazine 3 gr qd. Rapid remission of all symptoms was noted as well as decreased inflammatory parameters, including Ig G4.
Although EUS after 4 weeks of treatment was identical to the initial procedure; the biliary stent was removed and no cholestasis recurrence was noted.

At 5-month the patient is in complete remission without evidence of autoimmune pancreatic activity (i.e., without signs or symptoms of pancreatic insufficiency or cholestasis).

Discussion

The diagnosis of AIP is a clinical challenge, not only due to its rarity, but also due to the need of integrating clinical, laboratory, imaging and histology data for confirmation.6,7 Because of that, AIP patients are frequently submitted to multiple exams, and some of which are invasive, until a definitive diagnosis can be reached. The clinical case presented here is an example of that, much because of the absence of characteristic imaging (such as the lack of the "sausage-like" aspect of the pancreas on the CT or the identification of focal pancreatic lesions) and the inability to obtain a pancreatography by ERCP, which in case of AIP typically reveals focal segmental or diffuse stenosis, with little or no dilatation of the amount of segments.6-9 Therefore, EUS proved fundamental in this case. Although no imaging criteria can be considered pathognomonic, morphology on EUS raised the suspicion which lead to the decision of obtaining pancreatic tissue,8,10,11 underscoring the fact that histological examination by an experienced pathologist could be considered the gold standard.6,7,12-14

The association of AIP with other autoimmune illnesses can be identified in more than half of the cases.11,13 They can precede the pancreatic illness diagnosis or present later during the natural course of the disease.9 Among these, the association with IBD, and more specifically with UC, has been described, being the most common in an Italian series (35% of analysed cases).9,10,13 Overall, however, the true dimension of the relationship between these two entities is still not totally clear. This is likely due to the fact that only recently AIP has been considered a proper nosological entity with well defined diagnostic criteria. Indeed, we believe that previous case reports referring to chronic pancreatitis with indeterminate aetiology associated to UC probably were cases of autoimmune pancreatitis based on current diagnosis criteria.15

In fact, the observed pancreatic alterations of patients with UC are more frequent than initially expected. Although UC patients present an increase incidence of gallbladder lithiasis and are administered drugs that can potentially be pancreato-toxic, these factors alone are probably not enough to explain the great incidence of pancreatic alterations among UC patients.16 Some studies demonstrate insufficient levels of pancreatic exocrine in 21–80% of IBD patients and autopsy studies register pancreatic alterations, macroscopic or microscopic, in 14–53% of UC patients. Pancreatic duct changes, such as irregularities or short-segment stenosis of the main pancreatic duct, were observed in 8.4–10.8% of IBD patients independently of prior history of pancreatitis or exocrine insufficiency.4,17 It seems that predominantly asymptomatic pancreatic alterations of indolent development might exist in these patients, albeit the fact that the exact aetiology and pathogenesis are still poorly understood. We believe that a large spectrum of pancreatic changes can be documented in IBD patients, from symptom-free cases (likely the majority) to clinically exuberant forms such as the case of our patient. The aetopathogenesis could be related to an abnormal immunological response leading to pancreatic inflammation such as Ectors et al. previously suggested.18

The association between AIP and UC presents a clinical challenge concerning the treatment strategy. UC patients need immunosuppressive treatment in up to 30% of cases.19 Thiopurins (azathioprine and 6-mercaptopurine) continue to be the most widely used. However, potential pancreatic adverse effects are well established, raising concerns of its use in patients with AIP. In this setting other therapies (e.g. methotrexate or biological therapy) could step-in as first line options.20 There are some authors who advise against the use of thiopurins in AIP, although its use has been described as presenting good results in cases of relapse of AIP with a low level of adverse effects.21-23 Although more studies are needed, its use can be justified to avoid long-term treatment with corticosteroids, under close monitoring for pancreatic toxicity.

In respect to corticotherapy, a good clinical response is considered by some groups as a diagnostic criterion for AIP.7 In our case, a clinical and analytical improvement was seen, with no cholestasis relapse after biliary stent removal. Pancreatic morphology improvement on EUS was not observed after corticotherapy, supporting the idea of an irreversible extensive fibrotic process.11

A word of caution is in order, concerning the uneventful evolution of the presented case. A long-term follow-up strategy is mandatory, namely to maintain a low threshold for future associated autoimmune illnesses.

Conflicts of interest

The authors have no conflicts of interest to declare.

References