**Titel**  Autoimmune pancreatitis exhibiting multiple mass lesions

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**Short title**  AIP with multiple mass lesions

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Abstract  Our case is a first report of autoimmune pancreatitis (AIP) with multiple masses within the pancreas which was pathologically diagnosed by endoscopic US-guided fine-needle aspiration (EUS-FNA) and treated by steroid. The masses disappeared by steroid therapy. Our case is informative to know that AIP sometimes exhibits multiple masses within the pancreas and to diagnose it without unnecessary surgery.

Introduction

Autoimmune pancreatitis (AIP) is a unique form of chronic pancreatitis associated with an autoimmune inflammatory process [1, 2]. Although diffuse swelling of the pancreatic parenchyma and diffuse irregular narrowing of the pancreatic duct system are morphologically characteristic of AIP, a focal type of this clinical entity has been recently recognized [3]. The focal type of AIP exhibits a localized mass lesion in the pancreas, similar to pancreatic carcinoma [4]. Consequently, some patients with these features have been subjected to surgical exploration with a presumed diagnosis of pancreatic carcinoma. Considering that AIP shows a favorable response to steroid therapy, the differentiation of these two entities is clinically important to avoid surgery.

There have been only a few cases of AIP with multifocal lesions [5] but no case
pathologically diagnosed by EUS-FNA and treated by steroid. In this report, we describe the clinical, radiological, and histopathological features of a patient with AIP who exhibited distinct double masses in the pancreas and was treated by steroid.

Case report

A 63-year-old male patient without any symptoms was admitted because of pancreatic masses that were picked up on a medical checkup. He had a history of diabetes mellitus but none of alcohol abuse. On physical examination, the patient showed bilateral swelling of the submandibular gland. On laboratory examination, his serum glucose was 179mg/dL (normal range 65–109 mg/dL), γ-GTP was 132IU/L (normal 54.0 IU/L), cancer antigen 19-9 (CA19-9) level was 55.5 ng/mL (normal 37.0 ng/mL) and serum IgG4 was 773 mg/dL (normal range 4.8-105 mg/dL). Other serological tests, including pancreatic and hepatobiliary enzymes, γ-globulin, immunoglobulinG, and tumor markers (CEA and DUPAN-II) were within the normal range. Antinuclear antibodies (ANA) were also negative at a titer of less than 1 : 20.

On abdominal ultrasonography [6], the patient was found to have low echoic mass in the pancreatic head uncus and body. Contrast-enhanced computed tomography (CT) showed two mass lesions in the head and body of the pancreas (Fig. 1A, 1B); the lesions
were 18 mm and 10 mm in diameter, respectively. The tumors showed slight attenuation in the delayed phase (Fig. 1C, D). On dynamic magnetic resonance imaging (MRI), they were also hypovascular during the early phase and showed delayed enhancement during the late phase. Endoscopic retrograde cholangiopancreatography (ERCP) showed localized narrowing of the main pancreatic duct at the pancreatic head. The common bile duct was not dilated. Fluorine-18 fluorodeoxyglucose-positron emission tomography (FDG-PET) showed FDG uptake in the bilateral submandibular glands, hilar, mediastinal lymph nodes and pancreatic head uncus and no uptake in the body of pancreas.

EUS-FNA of the two mass and normal pancreas was performed and the specimen were adequate for cytology and cell block. Cytology in the two mass was rich in inflammatory cells. The cell block in the two mass revealed a dense lymphoplasmacytic infiltrate and significant replacement of pancreatic parenchyma by irregular fibrosis (Fig.2A). Immunostaining for IgG4 revealed diffuse infiltrate of IgG4-positive plasma cells (Fig.2B). Cytology and the cell block in the normal pancreas had no abnormality (Fig.2C).

On the basis of the above-mentioned clinical, imaging, and cytological findings, a diagnosis of autoimmune pancreatitis was made, and treated with prednisone, which
was initiated at a dose of 30 mg per day with a tapering schedule of 5 mg every 2 weeks. One month after treatment, the elevated IgG level resolved and CT revealed that the two mass disappeared (Fig. 3A, B, C) and FDG uptake in the pancreas, submandibular glands, hilar, and mediastinal lymph nodes disappeared completely.

**Discussion**

We showed the clinical, radiological, histopathological and clinical features of a patient with AIP who had distinct double masses in the pancreas and steroid therapy.

Since Yoshida et al. proposed the term “autoimmune pancreatitis” in 1995, this unique pancreatic disorder was recognized as a new clinical entity with such characteristics as increased serum γ-globulin, IgG levels, the presence of autoantibody, diffuse enlargement of the parenchyma, and irregular narrowing of the pancreatic duct [1, 7]. In 2001, Hamano et al. reported that patients with AIP have a high serum concentration of IgG4 [6]. The specificity and sensitivity of a high serum IgG4 level in the diagnosis of AIP are higher than 90% and this is now believed to be the most useful examination for diagnosing AIP. Hirano et al. [8] recently reported eight cases of AIP with locally affected lesions. They found that the localized type of AIP is more difficult
to differentiate from pancreatic cancer radiologically because typical radiological features reflecting diffuse involvement are not observed. They reported that serum IgG4 was elevated in the cases of focal AIP and that serum IgG4 levels might be helpful for its diagnosis. However, a pancreatic cancer with elevated serum IgG4 was reported [9].

Multiple masses in the pancreas are a rare clinical entity. As previously reported, multiple pancreatic masses were as follows: metastatic pancreatic tumors such as renal cell carcinoma[10], pancreatic neuroendocrine tumor (PNET) in multiple endocrine neoplasia type 1 (MEN1) [11], pancreatic cancer[12], and autoimmune pancreatitis (AIP) [5, 13]. Our patient had no primary lesion other than pancreas and the imaging findings in the present case were not suitable for PNET but pancreatic cancer could not be excluded based on the radiological images in the present case. Radiological findings were indicative of a malignant pancreatic tumor, especially pancreatic cancer. And also, few reports on the coexistence of AIP and PDAC have been published posing a new problem in the management of AIP [14, 15].

Endoscopic US-guided fine-needle aspiration of the pancreas or EUS trucut biopsy (TCB) are useful for diagnosing AIP and excluding pancreatic cancer [16]. FNA in this patient was compatible with AIP. However, pancreatic cancer is rarely difficult to exclude by biopsy specimens alone because of reactive inflammatory cell reactions and
fibrosis surrounding the neoplastic cells.

It is interesting in the present case that multiple mass were formed in the pancreas. These multiple lesions have been rarely reported for AIP until now. We could not explain this unusual manifestation, but early phase of AIP could be focal lesions.

The most important thing is that we recognize that AIP can manifest as multiple mass lesions similar to those in the present case, examine IgG4, perform EUS-FNA or TCB-FNA and try to deny pancreatic cancer. If IgG4 is elevated and EUS-FNA exclude malignancy, steroid therapy should be carefully challenged, quickly check the response by CT or PET, and the patient may avoid unnecessary surgery. This case is a first report of AIP with multifocal masses treated by steroid and we believe this diagnostic process is informative.


**Figure legend**

**Fig. 1A–D.** Computed tomography (CT) revealed mass lesions in the head (A) and body (B) of the pancreas (*arrows*): early phase. They showed slight delayed enhancement during the late phase (*arrows in C, D*).

960 × 720 pixel

**Fig. 2A, B, C** Cell block in the two nodule showed dense lymphoplasmacytic infiltrate and significant replacement of pancreatic parenchyma by irregular fibrosis (H&E×400) (A). Numerous plasma cells in the two nodule show positive immunoreactivity for IgG4 (H&E×400) (B) Cell block in the normal pancreas showed almost normal acinar cell, no lymphoplasmacytic infiltrate and no fibrosis (H&E×400) (C).

960 × 720 pixel

**Fig. 3A, B** The two nodule in the head(A) and body(B) of CT disappeared.

960 × 720 pixel