Pancreatic hamartoma, a rare benign disease of the pancreas:
A case report

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Abstract. Pancreatic hamartoma is an extremely rare, non-neoplastic, mass-forming lesion that may be mistaken for malignancy, and the pre-operative diagnosis is particularly challenging. The published literature contains only 23 cases of pancreatic hamartoma. The majority of the cases reported patients with a single benign tumor-like disease that received a pancreatectomy. Immunohistochemical findings confirmed the diagnosis following surgery. The current study reports the case of a 53-year-old female who presented to the Department of Abdominal Surgery, Chinese Academy of Medical Sciences, Cancer Hospital (Beijing, China), due to abdominal pain. Abdominal magnetic resonance imaging revealed a 22x14-mm mass in the head of the pancreas. The patient was pre-operatively diagnosed with a pancreatic space-occupying lesion, and subsequently underwent a pancreatectomy. The post-operative course was uneventful. Histological examination of the resected lesion resulted in a diagnosis of pancreatic hamartoma. There were no signs of recurrence at 55 months post-surgery.

Introduction

Hamartomas are benign tumors formed from the disordered growth of normal cells and tissues at the affected site (1). Pancreatic hamartoma is extremely rare and accounts for <1% of hamartoma; the etiology is unknown and the disease has a low morbidity (2). With the increased utilization of imaging technology, such as endoscopic ultrasound, there has been an increased detection of asymptomatic lesions of the pancreas; however, the differentiation between hamartomas and other benign tumors or malignancies is extremely difficult (3). The final diagnosis is usually made according to histopathological and immunohistochemical results (4).

Due to the dangers of biliary obstruction or hemorrhage, a pancreatectomy is necessary (5). No recurrence usually occurs following radical excision. Pancreatic hamartoma can be divided into two subgroups: Solid and cystic, or solid (6). The majority of cases reported in the literature are cystic hamartomas (1-3,7-13). The present study reports the case of a solid pancreatic hamartoma that was diagnosed after surgery.

Case report

A 53-year old female was admitted to the Department of Abdominal Surgery, Chinese Academy of Medical Sciences, Cancer Hospital (Beijing, China) in October 2014, due to a 2-month history of abdominal pain, accompanied by slight anorexia and 2 kg of weight loss. Physical examination did not detect any abnormalities. The history of the patient did not disclose any systemic diseases or previous surgeries. The patient reported no alcohol consumption or cigarette use, and there was no family history of tumor-associated disease. Blood tests indicated that tumor marker levels [carcinoembryonic antigen, carbohydrate antigen (CA)19-9, CA-125 and CA72-4] were within the normal ranges, and pancreatic exocrine and endocrine function was sufficient. Other standard blood tests were unremarkable. Abdominal magnetic resonance imaging (MRI; MAGNETOM Skyra, Siemens Healthcare, Erlangen, Germany) was performed and revealed a 22x14-mm mass in the head of the pancreas. The patient was pre-operatively diagnosed with a pancreatic space-occupying lesion, and subsequently underwent a pancreatectomy. Endosonography (TGF-UC260J; Olympus Corporation, Tokyo, Japan) was also performed and revealed a 2.83x2.27-cm, hypoechogenic lesion located in the head of the pancreas. Endoscopic ultrasound (EUS) fine-needle aspiration was not performed due to the problematic position of the lesion, which was particularly close to the superior mesenteric vein (Fig. 2). The combined imaging results were not enough to form a reliable diagnosis; therefore, a formal pancreatectoduodenectomy was performed. During the surgery, a 2.3x1.5x1.5-cm, firm mass with an intact capsule was identified, which appeared to be embedded in the parenchyma of the head of the pancreas (Fig. 3). On the resected surface of the tumor, a well-demarcated,
solid, homogeneously white-colored nodule was identified. The resected tissue was formalin (Beijing Saichi Biological Technology Co., Ltd., Beijing, China)-fixed, paraffin (Beijing Saichi Biological Technology Co., Ltd., Beijing, China)-embedded and cut into 4-µm sections prior to microscopic examination (CX23; Olympus Corporation), which demonstrated that the lesion was composed of disarranged ductal and acinar cells, embedded in a markedly fibrous stroma. Hematoxylin-eosin staining indicated that the lesion was positive for α-1-antitrypsin, α-1-antichymotrypsin and s100, with a Ki-67 score of <1%. Small amounts of disordered acinar cells and ductal epithelium were immunohistochemically positive for cluster of differentiation (CD)117, CD56, chromogranin A, progesterone receptor and synaptophysin, and negative for cytokeratin 19, p63 and vimentin. Thus, the histological diagnosis was confirmed as pancreatic hamartoma. The post-operative clinical course was stable without complications, and the patient was discharged following 15 days of hospitalization. At 55 months of follow-up, no recurrence was observed.

Discussion

A hamartoma is a focal, benign malformation that resembles a neoplasm in the tissue of its origin (14). This lesion is not a real tumor, and it grows at the same rate as its adjacent tissues. Hamartomas may develop in various areas of the body, and are typically observed in the lungs, heart, kidneys, spleen or other vascular organs. These lesions are commonly asymptomatic and may remain undetected unless identified incidentally during imaging analyses (14).

Pancreatic hamartoma is extremely rare. Only 23 cases, including the present case, have been reported in the literature (Table I), with the first case being described by Anthony et al in 1977 (7). The development of this disease may occur at any age, even in young children, however, the average age of occurrence is 40-60 years (median, 42.23 years) (4,5, 8-10,15-18). Morbidity is not equal between men and women, with a male to female ratio of 1.5:1. In contrast to hamartoma in other organs, the majority of pancreatic hamartoma cases are accompanied by pain, abdominal discomfort or certain vague symptoms, including dyspepsia and weight loss (11,19). Tumors are often located in the head of the pancreas, but signs of jaundice are rare.
Table I. Clinicopathological features of pancreatic tumors reported as pancreatic hamartoma in the literature (n=23).

| First author/s, year | Age, years | Gender | Size, cm | Surgery | Pancreatitis | Solid and cystic/solid | Refs. |
|----------------------|------------|--------|----------|---------|--------------|------------------------|-------|
| Anthony et al, 1977  | 46         | M      | 1.6      | LD      | No           | Solid and cystic        | (7)   |
|                      | 35         | M      | Multiple | LR      | Yes          | NR                     |       |
|                      | 58         | M      | 1.0      | autopsy | No           | NR                     |       |
| Noltenius and Colmant, 1977 | 52      | F      | Multiple | autopsy | Yes          | NR                     | (15)  |
| Burt et al, 1983     | 0.65       | F      | 11.5     | TP      | No           | Solid and cystic        | (8)   |
| Flaherty and Benjamin, 1992 | 1.67    | F      | 9.0      | LR      | No           | Solid and cystic        | (9)   |
| Izbicki et al, 1994  | 25         | M      | 10.6     | PD      | No           | Solid and cystic        | (10)  |
| Wu et al, 1998       | 39         | M      | 8.0      | PD      | Yes          | NR                     | (16)  |
| McFaul et al, 2004   | 39         | M      | 1.0      | PD      | Yes          | NR                     | (19)  |
| McFaul et al, 2004   | 62         | M      | 3.5      | PD      | Yes          | NR                     |       |
| Pauser et al, 2005   | 36         | F      | 7.0      | PD      | No           | Solid and cystic        | (12)  |
| Pauser et al, 2005   | 55         | F      | 3.0      | DP      | No           | Solid and cystic        |       |
| Pauser et al, 2005   | 51         | M      | 3.0      | LR      | No           | Solid                  |       |
| Nagata et al, 2007   | 54         | F      | 2.0      | DP      | No           | Solid                  |       |
| Nagata et al, 2007   | 58         | F      | 1.9      | DP      | No           | Solid                  |       |
| Thrall et al, 2008   | 3          | M      | 3.0      | PD      | No           | Solid and cystic        | (11)  |
| Sampelean et al, 2009| 46         | M      | 8.0      | PD      | NR           | Solid                  | (17)  |
| Durczynski et al, 2011| 69     | M      | 3.0      | LR      | No           | Solid                  | (3)   |
| Kawakami et al, 2012 | 78        | F      | 1.8      | PD      | No           | Solid                  | (6)   |
| Kim et al, 2012      | 52         | M      | 2.2      | PD      | No           | Solid and cystic        | (2)   |
| Sueyoshi et al, 2013 | 3          | M      | 3.0      | PD      | No           | Solid and cystic        | (18)  |
| Addeo et al, 2014    | 61         | M      | 2.4      | DP      | No           | Solid                  |       |
| Present study        | 48         | F      | 2.2      | PD      | No           | Solid                  |       |

M, male; F, female; PD, pancreaticoduodenectomy; DP, distal pancreatectomy; LR, local resection; TP, total pancreatectomy; NR, no record.
Pancreatic hamartoma is typically composed of disorganized, well-differentiated exocrine and endocrine pancreatic tissue. Acinar, islet and ductal cells are the three main components that form the lesions (1). A specific cell type, termed spindle-shaped cells, exhibit immunoreactivity for CD34 and CD117 in a number of pancreatic hamartoma cases (1,12). Pancreatic hamartoma is classified as solid or solid and cystic by macroscopic findings (13). Although the pathogenesis and origin of this disease remains unclear, it should only be diagnosed in patients without evidence of chronic pancreatitis, as chronic pancreatitis often presents with a depletion of acinar cells in the fibrous stroma, thus mimicking hamartoma with a lack of such cells (12).

To date, the existing imaging methods employed to characterize pancreatic hamartoma lack the accuracy to successfully distinguish it from other diseases. The majority of lesions that develop in the pancreas are malignant or have malignant potential, therefore, surgery is the optimal treatment in such instances, and a complete resection with negative margins is imperative (3). The treatment of choice for hamartoma is a conventional pancreatectomy or a pancreas-preserving surgery (central pancreatectomy or enucleation). A conventional pancreatectomy, including pancreaticoduodenectomy and distal pancreatectomy, may sacrifice the normal pancreatic parenchyma and be associated with the risk of post-operative diabetes mellitus or endocrine and exocrine pancreatic insufficiency (2). Considering the benign behavior of this tumor, pancreas-preserving surgery is recommended, which has the advantages of preserving the integrity of the gastrointestinal tract and splenic function, and sparing the maximal pancreatic endocrine and exocrine pancreatic insufficiency (2). In conclusion, pancreatic hamartoma may be detected incidentally, without the patient presenting with any signs or symptoms of disease. Clinical symptoms are dependent on tumor size and location, and the diagnosis of this disease primarily depends on imaging techniques, including computed tomography, MRI and EUS. However, pancreatic hamartoma is difficult to differentiate from other benign pancreatic lesions. For symptomatic patients or those with an indefinite diagnosis, a complete surgical excision is recommended. Following a successful pancreaticoduodenectomy, the patient in the present study was diagnosed with a pancreatic hamartoma on histological examination, and no recurrence has since been observed.

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