Serous cystadenoma of the pancreas with atypical clinical manifestations: A case report

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Abstract. The present study reported a 4.8 cm space-occupying cystic solid mass of the pancreas, which caused recurrent bilateral lower back discomfort in a 60-year-old female. Combined with the clinical data, an impression of a solid-pseudopapillary neoplasm was generated prior to surgery. Abdominal exploration revealed splenic vessel encasement by the mass similar to the invasion observed in a malignant tumor. Distal pancreatectomy with splenectomy was subsequently performed. Grossly, the tumor was solid and consisted of numerous small cysts. Histopathological examination of the cystic solid mass revealed classic microcystic serous cystadenoma of the pancreas. These findings suggested that microcystic serous cystadenoma can exhibit atypical clinical manifestations. Asymptomatic patients with a small lesion (<4 cm) require imaging surveillance every 2 years; however, tumors >4 cm with atypical presentations require surgical resection.

Introduction

Serous cystadenoma of the pancreas (SCA) is a benign epithelial neoplasm of uncertain origin. Immunohistochemical and ultrastructural studies have suggested that SCA may originate from centroacinar or intercalated duct cells. It can occur in any region of the pancreas, predominantly in older females (1). According to the literature, the majority of SCAs are small (<1 cm) and are identified incidentally. Grossly, SAC is a well-circumscribed mass within numerous small cysts (‘honeycomb’ appearance) on cut surface with a central fibrotic scar. Histologically, majority of SACs are ‘microcystic’ with numerous cysts exhibiting a diameter <2 cm and rarely with calcification in the fibrous septa. Lesions with a diameter >2 cm were defined as macrocystic serous cystadenoma (2,3).

Discussion

Serous cystadenoma of the pancreas was first described by Compagno and Oertel in 1978 (1). Morphological features
include numerous cysts with variable diameter lined by a single layer of cuboidal epithelium producing watery fluid similar to serum. SCA occurs most frequently in females ranging between 56 and 62 years of age (4, 5). SCA can occur in any section of the pancreas, although predominantly in the head and body region (6). SCA is associated with von Hippel-Lindau disease and can have diffuse and multiple lesions over the entire pancreas at a younger age in this setting (7). CT and MRI are the most common diagnostic methods for SCA. CT and MRI imaging of SCA typically reveals a lobulated contour, uneven low density, thin wall/capsule and absence of wall enhancement. Atypical imaging features include multiple septations, wall enhancement and/or calcification, wall thickness, and presence of mural nodules or papillary projections (2). However, these features cannot differentiate SCA from other pancreatic cystic tumors. Approximately 60% of the SCAs can be diagnosed by CT and/or MRI. When imaging features reported by CT and/or MRI are insufficient to differentiate cystic lesions, endoscopic ultrasonography (EUS) with or without EUS-guided fine-needle aspiration (FNA) can provide additional and useful information, and notably improves the sensitivity of diagnosis for SCA to 76% (8). Although EUS with FNA increases the detection rate of SCA, FNA is somehow limited due to a high rate of insufficient sampling and potential tumor seeding of the needle tract for cases when malignancy cannot be ruled out. Serum CEA and CA19.9 are non-specific biomarkers for pancreatic lesions.

Almost all SCAs are benign and to date, no patient has succumbed to SCA or associated reasons (9, 10). Therefore, surgical resection for serous cystic neoplasms should be limited to those with clinical symptoms and large tumor size. A previous study indicated that a tumor > 6 cm is the cornerstone for operation (11). Based on our experience, the present study suggested that asymptomatic patients with a small lesion (< 4 cm) require imaging surveillance every 2 years. However, tumors > 4 cm require surgical resection, since these tumors are more frequently symptomatic and faster growing (12). When multiphasic contrast enhanced CT scan and MRI imaging studies reveal a low diagnostic power to differentiate SCA from mucinous cystic neoplasm and intraductal papillary mucin neoplasm, early surgical treatment must be taken into account with growing experience and advanced surgical techniques.

Surgical resection is considered curative for SCA. The location of SCA in the pancreas determines the method of resection, which is used. The appropriate methods for this case include distal pancreatectomy with spleen-preserving or with splenectomy. Spleen-preserving distal pancreatectomy includes the Warshaw procedure, which conserves the spleen by blood flow from the short gastric vessels and the Kimura procedure, which preserves the spleen with splenic vessels (13). Since the Warshaw procedure has a high incidence of complications and subsequent splenectomies, distal pancreatectomy with splenectomy is often required, particularly when the splenic vessels are encased by SCA, as in this
case. Therefore, when no recurrence is observed following radical surgery, post-operation follow-up is unnecessary in these patients.

In the previous decade, non-surgical management for patients with SCA is rising. EUS-guided treatment has been reported, particularly for patients unfit for surgery. Cyst ablation with the injection of ethanol followed by paclitaxel was considered as another promising technique and has revealed complete resolution in certain patients, although long-term follow-up is still required (14,15).

In conclusion, with the most recent imaging techniques, a large proportion of patients with SCA cannot be definitively diagnosed prior to operation, particularly with atypical imaging findings. Although SCA is a benign pancreatic cystic neoplasm, the patient can be symptomatic, particularly when the cyst is >4 cm and surgical treatment is particularly important to manage these patients.

References