Heterotopic pancreas in the stomach: A case report and literature review

Grigoris Christodoulidis, Dimitris Zacharoulis, Sotiris Barbanis, Emmanuel Katsogridakis, Konstantine Hatzitheofilou

INTRODUCTION

Pancreatic heterotopia was first described in 1727 when it was found in an ileal diverticulum[1]. It is a rare entity, defined as the presence of extrahepatic tissue without any anatomic or vascular continuity with the pancreas. It may occur at a variety of sites in the gastrointestinal tract having a propensity to affect the stomach and small intestine. Usually, it is a silent anomaly but it may become clinically evident when complicated by inflammation, bleeding, obstruction or malignant transformation[2]. We report a case of a 40-year-old female with an ectopic pancreatic lesion in the antrum of the stomach.

CASE REPORT

A 40-year-old woman was admitted to our hospital due to a 2-mo history of recurrent episodes of epigastric pain, nausea and vomiting. Physical examination, routine blood tests including amylase, plain chest and abdominal X rays along with abdominal ultrasound were unremarkable. Esophagogastroduodenoscopy revealed a sessile polypoid mass with benign features located in the gastric antrum to the posterior wall measuring approximately 2 cm in diameter. The mucosa appeared normal throughout the stomach. Biopsy confirmed the presence of a normal gastric mucosa over the lesion. Computed tomography was not performed.

A decision was made to proceed with surgery. Endoscopic injection with methylene blue was performed to mark the lesion preoperatively.

The patient underwent exploratory laparotomy. Through a small midline incision a gastrotomy was performed. The lesion was clearly stained with methylene blue 4 h after the endoscopy. It was located approximately 10 cm from the pylorus to the greater curvature. In palpation the lesion was rubber like, fixed to the surrounding mucosa giving the feeling 'like a breast fibroadenoma'. Its dimension was approximately 5 cm × 3 cm × 4 cm. By using a stapler device a wedge resection of the lesion was performed with macroscopically clear margins. Frozen sections excluded malignancy and the possibility of ectopic pancreatic lesion. The surgical margins were clear. The patient had no postoperative complications and was discharged 4 d later. She has remained free of symptoms with negative endoscopy since then.

Histopathologic examination of the lesion showed heterotopic pancreatic tissue in the gastric antrum with a lobular architecture characteristic of ectopic pancreas. The pancreatic lobules were located mainly in the gastric submucosa (Figure 1A) and partially in the muscularis propria (Figure 1B). They contained a mixture of pancreatic acini, ducts and islets of Langerhans. The overlying gastric mucosa was normal.

DISCUSSION

Ectopic pancreas is defined as pancreatic tissue that lacks anatomical or vascular communication with the normal...
Heterotopic pancreas has a genetic make-up, physiologic function, and local environmental exposure similar to that of the pancreas\[^5\]. The incidence in autopsies ranges 0.5%-13.7\%, being more common at the age of 30-50 years with a male predominance\[^3\].

Of the 105 gastrectomies performed in our institution over the last five years, ectopic pancreatic tissue was found in only one case (1/105, 0.9\%).

Several theories have been proposed to explain the pathogenesis and occurrence of pancreatic heterotopia. The most tenable theory implicates that during the development of normal pancreas from several evaginations, originating from the wall of the primitive duodenum, one or more evaginations may remain in the bowel wall. Migration of this embryonic remnant along with the development of the gastrointestinal tract gives rise to the ectopic pancreatic tissue\[^5\]. Another theory suggests that during embryogenesis pancreatic metaplasia of the endodermal tissues localized in the gastric submucosa may occur\[^5\].

Histopathologically, it is not a diagnostic problem when pancreatic acini, ducts, islets of Langerhans and intervening connective tissue are present. The most characteristic gross feature is a central ductal orifice\[^6\].

Specifically in the stomach, the involvement of submucosal layer, muscularis and subserosal layer is 73\%, 17\% and 10\%, respectively\[^7\]. In the presented case the pancreatic tissue involved both the submucosa and muscularis propria. Heinrich in 1909 proposed three types of heterotopic pancreas but his classification was modified by Gaspar-Fuentes in 1973 acquiring its final form. Type I heterotopia consists of typical pancreatic tissue with acini, ducts, and islet cells similar to those seen in normal pancreas (Figure 1). Type II heterotopia is composed of pancreatic ducts only, referred as canalicular variety. Type III heterotopia is characterized by acinar tissue only (exocrine pancreas). Type IV heterotopia is made up of islet cells only (endocrine pancreas)\[^8\].

The usual location is in the stomach in 25%-38\% of the cases, duodenum in 17%-36\%, jejunum in 15%-21.7\% and rare in the esophagus, gallbladder, common bile duct, spleen, mesentery, mediastinum and fallopian tubes. Gastric lesions are discovered in the antrum in 85%-95\%, either on the posterior or anterior wall, being more common along the greater curvature\[^9\].

The pancreatic ectopic tissue is usually silent but can also undergo complications that occur in normal pancreatic tissue such as acute or chronic pancreatitis, abscess and pseudocyst formation\[^9\]. Malignant transformation may rarely occur. Up to 15 cases have been reported so far\[^10\]. In order to be described as arising from heterotopic pancreas, the diagnosis of a carcinoma should fulfill three criteria: (1) the tumour must be located within or very close to the ectopic pancreatic tissue, (2) transition between pancreatic structures and carcinoma must be identified and (3) the non-neoplastic pancreatic tissue must comprise fully developed acini and ducts\[^11\]. Adenocarcinomas arising from ectopic pancreas seem to have a somewhat better prognosis than those arising from the pancreas itself, probably due to earlier presentation\[^10\].

Symptoms depending upon the anatomical location, such as gastric outlet obstruction in a pre-pyloric rest or obstructive jaundice in a bile duct focus, may originate from the mass effect of the tumour\[^12\] and are also related to the size of the lesion. Lesions greater than 1.5 cm in diameter are more likely to cause symptoms\[^13\]. Pain is one of the most common symptoms. The possible explanation is that the pain is due to endocrine and exocrine function of the heterotopic pancreatic tissue, and relates to the secretion of hormones and enzymes, being responsible for inflammation or chemical irritation of the involved tissues\[^14\]. Haemorrhage due to mucosal erosion, ulcer formation and perforation especially localized in the small intestine have also been reported\[^15\].

Barium swallow study may show a typical image of a rounded filling defect with central indentation. The reported sensitivity and specificity are 87.5\% and 71.4\%, respectively\[^14\]. Upper GI endoscopy can demonstrate a broad based umbilicated submucosal lesion. In the majority of cases, biopsies are superficial and non diagnostic. However, positive biopsies can establish the diagnosis\[^15\]. Endoscopic ultrasonography has proven to be a useful adjunct in identification of pancreatic rests, localizing in the submucosa and ranging 0.5-2 cm. The combination of endoscopic ultrasonography with fine-needle aspiration allows cytologic evaluation of submucosal gastrointestinal lesions, having a sensitivity ranging 80%-100\%\[^16,17\].

Computed tomography findings are usually non specific. However, multi-slice spiral CT with oral and portovenous phase IV contrast may demonstrate the lesion which enhances similarly with the normal pancreatic tissue. CT can localize lesions with normal pancreatic tissue but cannot distinguish ectopic pancreas from other submucosal tumors\[^18,19\].

In our case, since neither CT nor endoscopic ultra-
sonography was performed and the biopsy showed normal gastric mucosa, the diagnosis was made based on the benign endoscopic features of the lesion.

The diagnosis may be sometimes difficult intraoperatively due to the gross similarity of pancreatic heterotopia with gastrointestinal stromal tumour (GIST), gastrointestinal autonomic nerve tumour (GANT), carcinoid, lymphoma or even gastric carcinoma. If in doubt, frozen section can help to avoid unnecessary extensive operations.

In conclusion, although pancreatic heterotopia is rare, it should be always considered in the differential diagnosis of extramucosal gastric lesions. Despite the development of modern diagnostic modalities, its diagnosis remains challenging. Surgical excision provides symptomatic relief and is recommended especially if diagnostic uncertainty remains. If in doubt, frozen section can help to avoid unnecessary radical operations.

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