Symptomatic pancreatic heterotopia treated by local excision

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Abstract

Non-ulcer dyspepsia is a continuing problem and in many cases a precise cause is never identified. We present five patients with an allegedly uncommon condition—pancreatic heterotopia. They were managed by local excision of the tumour and after a mean (range) follow up of 42 (9–80) months all remain free of the original symptoms.

Patients and methods

The details of the five patients are summarised in Table. It can be seen from this, that their presenting symptoms were both variable and non-specific.

All five patients presented with upper gastrointestinal symptoms; three complaining predominantly of dyspepsia and two with a history suggestive of gastric outlet obstruction, associated in one patient with a succession splash. Three of the patients also complained of alteration in bowel habit, which in two resolved after operation.

The antral tumour was found on gastroscopy in all five patients. In each case the appearance was of a small, smooth extra mucosal lump, which in two instances had a central punctum. Endoscopic biopsy specimen, however, showed pancreatic heterotopia in only two patients. All five patients had histological evidence of chronic gastritis, which was associated with the presence of Helicobacter pylori in one. Barium meal examination outlined the lesion in two of the four patients in whom it was used. The radiological findings of a small intramural mass with a central barium filled niche (Fig 1) were as described in other reports of this condition.

The five patients all underwent laparotomy and local excision of the tumour, which in each case was located within 3 cm of the pylorus (Fig 2). In one patient the excision was closed as a pyloroplasty, owing to the very close proximity of the lesion to the pylorus. Histology of the excised specimen showed heterotopic pancreas in each case. All the patients made an uneventful postoperative recovery. At a mean (range) follow up of 42 (9–80) months all the patients remain free of their original symptoms.

Discussion

Pancreatic heterotopia is defined as the presence, outside its usual location, of pancreatic tissue which lacks anatomical and vascular continuity with the pancreas proper. It has a reported incidence of 1–14% on necropsy examination and has been observed in 1 in 500 upper abdominal operations. Pancreatic heterotopia accounts for up to 7% of all benign gastric tumours.

In a series of 37 patients reported by Lai and Tompkins, the heterotopic pancreas was judged...
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Figure 2: Operative photograph from patient five showing heterotopic pancreatic tissue in gastric antrum with punctum cannulated by probe.

Symptomatic pancreatic heterotopia was treated by cannulation. Figure 2 shows the heterotopic pancreatic tissue in the gastric antrum, as in the five patients reported here.

The role of pancreatic heterotopia in producing symptoms is not clear – overlying gastritis, pylorospasm, and gastric outlet obstruction by larger tumours have all been suggested as possible mechanisms. The symptoms may not be specific for pancreatic heterotopia, however, as other benign prepyloric tumours can present in a similar manner.

Lower gastrointestinal symptoms have not been a feature in previous reports of pancreatic heterotopia. The three patients in this study who complained of alteration in bowel habit were all in the younger age group and coincidental causes, particularly the irritable bowel syndrome, cannot be ruled out. It is of note, however, that the altered bowel habit resolved in two of the three patients immediately after operation, which suggests a true association. It is interesting that these were the same two patients who had clinical findings suggestive of gastric outlet obstruction, which has frequently been found to be associated with altered bowel habit, following the original report by Dragstedt et al in 1947.

Symptomatic patients require surgical exploration in order to obtain a definitive diagnosis and to exclude malignancy. Local excision is adequate if the lesion looks benign. This proved to be safe and effective in the treatment of the five patients reported here. Although any operation may be associated with a placebo effect, we feel that the relief of symptoms in our five patients resulted from the removal of a pathological lesion.

We conclude that pancreatic heterotopia may be responsible for gastrointestinal symptoms. Although uncommon, we feel that it is an important diagnosis to consider in patients found to have a benign looking tumour in the stomach or duodenum on gastroscopy or barium meal examination since it can be simply and effectively managed by local excision.