Heterotopic Pancreas Presenting With Intussusception - A Rare Case Report

Sainath K. Andola*¹, Shilpa I. Bhimalli², Anita A. Mahanta³ and S.R. Harwal⁴

¹Professor and HOD, Department of Pathology, Mahadevappa Rampure Medical College, Gulbarga, India
²Resident Department of Pathology, Mahadevappa Rampure Medical College, Gulbarga, India
³Associate Professor Department of Pathology, Mahadevappa Rampure Medical College, Gulbarga, India
⁴Professor Department of Pediatric Surgery, Mahadevappa Rampure Medical College, Gulbarga, India

*Corresponding author e-mail: drskandola@gmail.com

A B S T R A C T

A case of 14yr old boy with an ectopic pancreatic lesion in the ileal segment presenting with intussusception is being described. Histological examination revealed ectopic pancreatic tissue in submucosa of ileum with intussusception. The unusual site and difficulty of making an accurate diagnosis is highlighted with review of literature.

Keywords: Heterotopic pancreas, Intussusception, Ectopic pancreas.

INTRODUCTION

Pancreatic heterotopia was first described in 1727 when it was found in an ileal diverticulum¹ it is a rare entity, defined as the presence of extrahepatic tissue without any anatomic or vascular continuity with the pancreas. The term was first used by deCastro et al². 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 transformation³. A case of 14yr male with an ectopic pancreatic lesion in the ileal segment presenting with intussusception is being presented because of its unusual site.

CASE HISTORY

14 yrs, boy c/o fever since 1 week associated with severe pain abdomen since 1 day ultrasonography showed features of intussusception measuring around 8x4 cms.
All routine blood & urine examinations were done and are within normal limits.

Grossly Segment of ileum received measuring 26 cms in length, central part of wall showed blackish discoloration. One of surgical margins showed a nodular lesion measuring 2x1.5cms (fig. 1). Histological Section from intestine showed large areas of haemorrhage and acute inflammatory reaction with hyperplasia of Payer’s patches. Nodular mass revealed lobular structures consisting of round to oval cells in glandular and acinar pattern separated by thin fibrous septae. The nuclei were round with bland chromatin (fig. 2). Also seen were intralobular ducts. A diagnosis of heterotopic pancreas with intussusception was made.

**DISCUSSION**

Isolated heterotopic pancreas in the ileum is very rare and usually asymptomatic. It is found in about 1-2% of autopsies. Despite its congenital origin, pancreatic heterotopia was usually discovered in adults, being more common at the age of 30-50 years with a male predominance a sex ratio of 3:1. 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. The incidence in pediatric age group, varies from 6 to 16 %. In literature less than 25 cases of isolated heterotopic pancreas of ileum as leading point of intussusception in children being described. The relative frequency of a pathological lesion causing intussusception increases with age. However, the highest frequency of detection of pathologic lesion in cases. Of intussusception is still the first year of life. The present case is described in a 14yr old boy who presented with pain abdomen due to intussusception.

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 along with the development of the gastrointestinal tract gives rise to the ectopic pancreatic tissue. Histopathologically, the heterotopic pancreatic tissue may have all the elements of the normal pancreas which include pancreatic acini, pancreatic ducts, and islets of Langerhans.

Heinrich in 1909 proposed three types of heterotopic pancreas but his classification was modified by Gaspar-Fuentes in 1973 acquiring its final form. Type1 heterotopia is composed of pancreatic ducts only, referred as canalicular variety. Type 2 heterotopia is characterized by acinar tissue only (exocrine pancreas). Type 3 heterotopia is made up of islet cells only (endocrine pancreas). 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. Malignant transformation may rarely occur.

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. Barium swallow study may show a typical image of a rounded filling defect with central indentation. The lesion within the wall of the ileum may act as a lead point. This is thought to be the mechanism of intussusception. It has also been postulated that intussusception arises from local disturbance in the motility of the small intestine caused by the heterotopic
pancreas. The reported sensitivity and specificity are 87.5% and 71.4%, respectively.

Endoscopic ultrasonography has proven to be a useful adjunct in identification of pancreatic rests. CT findings are usually non specific. CT scan localize lesions with normal pancreatic tissue but cannot distinguish ectopic pancreas from other submucosal tumors. The diagnosis may be sometimes difficult intraoperatively due to the gross similarity of pancreatic heterotopias with gastrointestinal stromal tumour (GIST), gastrointestinal autonomic nerve tumour (GANT), carcinoid, lymphoma or even gastric carcinoma. If in doubt, frozen section is very helpful to establish the diagnosis intraoperatively and to avoid unnecessary extensive operations. In present case ultrasonoud showed signs of ileal intussusception. Exploratory laparotomy done and segment of ileum resected.

CONCLUSION

Isolated heterotopic pancreas in the ileum is very rare and usually asymptomatic. The lesion within the wall of the ileum may act as a lead point. This is thought to be the mechanism of intussusception. It should always be considered in the differential diagnosis of extra mucosal gastric lesions. Despite the development of modern diagnostic modalities, diagnosis remains challenging. Surgical excision provides symptomatic relief and is recommended if diagnostic uncertainty remains.

KEY MESSAGES

Ileal heterotopic pancreas is a rare cause of intussusception at any age it is often an incidental finding and is usually not evident clinically. Hence, careful examination of the lead point of intussusceptum in resection specimens is mandatory to delineate the underlying etiology of these cases.

REFERENCES

10. Grigoroius christodoulis, dimitris zacharoulis; Heterotopic pancreas in the stomach: A case report and literature review, world journal of gastroenterology; 2007; Dec 7; 13 (45).

**Figure 1.** Central part of ileum segment showing blackish discoloration

**Figure 2.** Microscopy reveals congested mucosal layer beneath which is seen ectopic pancreatic tissue in lobules (H & E 100x)