Case report on Ansa Pancreatrica: An Uncommon Cause Accounting for Recurrent Pancreatitis in Children
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Abstract
Acute recurrent pancreatitis is a rare entity in children. It can be caused by a number of reasons, anatomical variations being one of them. Pancreatica divisum is the most common form of ductal anomaly while ansa pancreatica has been the least studied and explored. In recurrent pancreatitis, Ansa Pancreatica was recently found to be a key risk factor. It is usually found among adult alcoholics. We submit the report of a rare but important cause of acute recurrent pancreatitis; an 11-year-old child with 2 previous episodes of pancreatitis was diagnosed with ansa pancreatica on magnetic resonance cholangiopancreatography (MRCP). He was advised to get stenting of Pancreatic duct. To the best of our belief, only another case has been reported in the paediatric population.

Keywords: Acute recurrent pancreatitis, ansa pancreatica, pancreatica divisum.

Introduction
Acute recurrent pancreatitis in childhood and adolescence is a rare occurrence and poorly understood. Few single-center studies have approximated 9-35% of children with acute pancreatitis experiencing recurrent episodes.1-3 The etiology in children differs from adults and comprises of genetic mutations, trauma, infection, drugs, anatomic malformations and metabolic disorders. Acute pancreatitis has a number of anatomical causes, of which pancreatica divisum is the most probable.4 Most anomalies and variants of pancreas are found incidentally and have no clinical significance. However, a few play a role in causing pancreatitis.

Ansa pancreatica, a rare ductal anomaly, bridges the main pancreatic and accessory pancreatic duct. In this pancreatic duct variant, a branch descends as an extension of the main pancreatic duct; later ascending to form a loop which terminates at the minor papilla. It is associated with recurrent acute pancreatitis due to causing hindrance in the drainage of pancreatic juice.5 Reports have shown that patients with ansa pancreatica, especially alcoholics, are likely to develop pancreatitis.4,6 During the literature review, despite there being a couple of case reports on ansa pancreatica in adults, there was only one reported in children.7

We submit a report of acute recurrent pancreatitis in an 11-year-old male, who on magnetic resonance cholangiopancreatography (MRCP) was discovered to have ansa pancreatica. The purpose of this case is to identify ansa pancreatica as a cause of recurrent pancreatitis in children.

Informed consent from the patient's parents and approval of the IRB committee was obtained for this study.

Case Report
An 11-year-old male child presented with severe epigastric pain, radiating to the back since one day on 13th September 2018. Concomitantly, he developed vomiting and nausea but no fever. He had a history of two similar episodes in June and August 2018 and following investigations had been diagnosed as acute recurrent pancreatitis. His family history for pancreatitis or any other disease was insignificant.

On admission, his abdomen was found to be non-tender and soft on examination without any visceromegaly. Normoactive bowel sounds were present on auscultation. Neither oedema nor palpable lymphadenopathy was present. General examination was unremarkable.

Patient was given supportive care and detailed laboratory workup was started. His total leukocyte count along with haemoglobin and liver function tests were normal. At admission, his serum amylase level was 523 International Unit per liter (IU/L) (Reference range 0-160 IU/L) and serum lipase was 1972 IU/L (Reference range: 1-130 IU/L). Serum calcium, sodium alongside triglycerides were within

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normal bounds. Patient underwent an extended autoimmune profile which was negative. An ultrasound of the abdomen revealed mild left renal pelvicalyceal fullness but it did not reveal any gallbladder stone or sludge. The patient was hospitalized and kept nil per os (NPO) while being managed with intravenous fluids and antibiotics.

His MRCP was performed which revealed a bulky pancreas with mild peripancreatic stranding consistent with findings of pancreatitis. It also showed mild intra and extrahepatic biliary dilation with narrowing of a small segment in distal common hepatic duct. Variable pancreatic ductal anatomy was noted showing the ventral duct (Wirsung duct) with no communication with the dorsal duct while draining into the minor papilla. The pancreaticogram revealed an arch ending adjacent to the minor papilla formed by a branch emerging from the main pancreatic duct [Figure-1]. This curved variant matches with the description of ansa pancreatica [Figure-2].

Following the antibiotics, the abdominal pain subsided and the patient’s condition improved in the next 5 days. Repeat serum lipase showed a considerable decrease to 1195 IU/L. He was discharged on low fat diet and regular follow-ups. Hepatobiliary consultation was given to the family with stenting and sphincterectomy recommended. The patient is currently doing well after sphincterectomy.

Discussion
Despite acute pancreatitis occurring more frequently in adults, there has been a recent rising trend in children. It carries a significant mortality and morbidity risk in children with increased economic burden requiring a mean stay of 13.2 days in the hospital and 28% of patients needing total parenteral nutrition. The known associations in children are genetic mutations, trauma, structural abnormalities, drug, viral infections while 23% are classified as idiopathic.

According to a study in Japan, amongst adult patients with acute pancreatitis, 2.9% had pancreatica divisum, 0.85% had ansa pancreatica while a winding main pancreatic duct was found in 6.3%. The risk of pancreatitis both acute and recurrent came out to be considerably higher in patients suffering from ansa pancreatica (20%) in contrast to the ones without it (0.52%).

Pancreatica divisum is the most frequent structural abnormality responsible for 5-8% of cases. One of the rarest cause is ansa pancreatica, where an arched duct replaces the normal accessory duct in the pancreatic head between dorsal and ventral ducts. This accessory pancreatic duct, with a reverse S-shape, branches from the main pancreatic duct, running into the caudal side of the ventral, terminating within or adjoining the minor papilla.

MRCP and Endoscopic retrograde cholangiopancreatography (ERCP) are considered to be the preferred diagnostic tests to identify the anatomical variant when it becomes symptomatic. In ERCP, the sigmoid branch of
ansa pancreatica may be wrongly identified as annular pancreas. However, differentiation can be done on the basis of a pancreatogram. In annular pancreas, the looping branch when approaching the minor papilla can be seen crossing the duodenum whereas in ansa pancreatica the branch does not cross the duodenum.

There is poor drainage of pancreatic juice in ansa pancreatica since the main pancreatic duct and side branch meet at a sloping angle, making patients vulnerable to pancreatitis especially in alcoholism and functional stenosis of sphincter of Oddi. This makes it a disease of the elderly.

However, the child in this case with relapsing episodes of acute pancreatitis proved ansa pancreatica to be a rare but possible differential diagnosis in the paediatric age group. Despite having sufficient data on other ductal and anatomical variations, ansa pancreatica has been largely ignored. Considering the paucity of data and lack of statistics, the need to follow outcomes and data becomes necessary for etiological assessment of recurrent pancreatitis and to give children a better quality of life.

**Conclusion**

Based on our case report, we conclude that despite being rare ansa pancreatica should be considered as a cause of recurrent pancreatitis in the paediatric age group. With surgical treatment, this disease could greatly impact the patient’s quality of life and should always be recommended.

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**References**
