Congenital Variants and Anomalies of the Pancreas and Pancreatic Duct: Imaging by Magnetic Resonance Cholangiopancreaticography and Multidetector Computed Tomography

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INTRODUCTION

Congenital anomalies and normal variants of the pancreas and pancreatic duct may not be detected until adulthood and they are often discovered as an incidental finding in asymptomatic patients. In adult patients with persistent and unexplained signs and symptoms such as recurrent abdominal pain, nausea and vomiting, recognition of these anomalies is important because these anomalies may be a surgically correctable cause of recurrent pancreatitis or the cause of gastric outlet obstruction. An awareness of these anomalies may help in surgical planning and prevent inadvertent ductal injury. The purpose of this article is to review normal pancreatic embryology, the appearance of ductal anatomic variants and developmental anomalies of the pancreas, with emphasis on magnetic resonance cholangiopancreatography and multidetector computed tomography.

Index terms: Pancreas; Congenital; Pancreas divisum; Annular pancreas; Accessory pancreatic lobe

Though congenital anomalies of the pancreas and pancreatic duct are relatively uncommon and they are often discovered as an incidental finding in asymptomatic patients, some of these anomalies may lead to various clinical symptoms such as recurrent abdominal pain, nausea and vomiting. Recognition of these anomalies is important because these anomalies may be a surgically correctable cause of recurrent pancreatitis or the cause of gastric outlet obstruction. An awareness of these anomalies may help in surgical planning and prevent inadvertent ductal injury. The purpose of this article is to review normal pancreatic embryology, the appearance of ductal anatomic variants and developmental anomalies of the pancreas, with emphasis on magnetic resonance cholangiopancreatography and multidetector computed tomography.
absent duct of Santorini and ansa pancreatica), duplication anomalies, anomalous pancreaticobiliary ductal junction, pancreas divisum, annular pancreas, ectopic pancreas, pancreatic agenesis and hypoplasia of the dorsal pancreas and accessory pancreatic lobe. Recognition of these anomalies is important because these anomalies may be a surgically correctable cause of recurrent pancreatitis or the cause of gastric outlet obstruction. An awareness of these anomalies may help in surgical planning and prevention of inadvertent ductal injury.

In this article, we review normal pancreatic embryology, the appearance of ductal anatomic variants and developmental anomalies of the pancreas, with emphasis on MRCP and multidetector computed tomography (MDCT).

**Embryologic Development of the Pancreas**

The pancreas develops from dorsal and ventral buds that first appear in the fifth gestational week as outgrowths of the primitive foregut (Fig. 1). By the seventh gestational week, expansion of the duodenum causes the ventral bud to rotate and pass behind the duodenum from right to left and fuse with the dorsal bud. The ventral bud forms the posterior head and uncinate process, whereas the dorsal bud forms the anterior head, body, and tail (3). Following this fusion, the ductal systems anastomose, a complicated process with a wide spectrum of possible outcomes. The portion of the ventral duct between the dorsal-ventral fusion and major papilla is termed the duct of Wirsung. The portion of the dorsal duct upstream to the dorsal-ventral fusion point is called the main pancreatic duct. The segment of the dorsal duct downstream to the dorsal-ventral fusion point is termed the duct of Santorini, or accessory pancreatic duct, which drains at the papilla minor (3).

**Variations of the Pancreatic Duct**

The course of the pancreatic duct varies greatly and the most common one (50%) is a descending course. Other courses include sigmoid, vertical, and loop configurations (Figs. 2, 3) (1).

The ductal configuration most commonly (60%) manifests as a bifid configuration with a dominant duct of Wirsung (Fig. 4). Less common configurations include an absent duct of Santorini (30%), a dominant duct of Santorini without divisum (1%), and ‘ansa pancreatica’, in which the duct of

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![Fig. 1. Drawings show normal embryologic development of pancreas.](image1)

![Fig. 2. Drawings show variation in course of pancreatic duct.](image2)

(A) Descending (B) vertical (C) sigmoid (D) loop shaped course.
Santorini forms a reversed-S shape and connects with a side branch of the duct of Wirsung (Fig. 5) (4).

Duplication anomalies of the main pancreatic duct are not uncommonly seen (Fig. 6). Cystic dilatations of terminal portions of the ducts of Wirsung and Santorini are termed Wirsungocele and Santorinicele (Fig. 7) (4).

Anomalous pancreaticobiliary ductal junction is characterized by fusion of the pancreatic duct and common biliary duct (CBD) outside the duodenal wall, with formation of a long common channel (> 15 mm) (Fig. 8). This anomaly
is often associated with choledochal cyst formation and carcinoma in the biliary tract (5).

**Developmental Anomalies of the Pancreas**

**Pancreas Divisum**

Pancreas divisum occurs in 4-14% of the population and results from failed fusion of the dorsal and ventral ducts during embryological development (3). Three variants have been described: type 1 or classical divisum in which there is total failure of fusion; type 2 in which dorsal drainage is dominant in the absence of the duct of Wirsung; and type 3 or incomplete divisum where a small communicating branch is present (6, 7). In pancreas divisum, the majority of the pancreatic gland drains into the minor papilla via the duct of Santorini, whereas the posterior head and uncinate process drain into the major papilla via the duct of Wirsung with the CBD (Fig. 9) (3). Although most patients are asymptomatic, in some patients, this anomaly is associated with recurrent acute pancreatitis because of inadequate drainage of pancreatic secretions via the minor papilla. MRCP enables a noninvasive diagnosis of the pancreas divisum without the use of contrast material and avoids the risk of endoscopic retrograde cholangiopancreaticography (ERCP) induced acute pancreatitis (1). MDCT may also depict this anomaly, when the pancreatic duct is visualized. On axial images, the dorsal duct passes the terminal common bile duct anteriorly and superiorly (Fig. 10) (8). Rarely, pancreas divisum is associated with a focal cystic dilation of the terminal portions of the duct of Santorini, which is termed as a Santorinicele (Fig. 7) (4).

**Annular Pancreas**

Annular pancreas occurs in 1/20000 of the population and results from failure of the ventral bud to rotate with the duodenum, resulting in envelopment of the duodenum. In this anomaly, a band of pancreatic tissue encircles the second part of the duodenum, either completely or incompletely, and it is in continuity with the pancreatic head (3). Annular pancreas usually presents in neonates with vomiting due to severe duodenal obstruction. In adults,
approximately 50% of the patients may be asymptomatic for life, with the anomaly discovered incidentally. When symptomatic, the presentation is usually in the third to sixth decade with abdominal pain, postprandial fullness and vomiting resulting from gastric outlet obstruction, upper gastrointestinal bleeding from peptic ulceration, acute or chronic pancreatitis, or in rare instances, jaundice resulting from biliary obstruction (9).

When the annular pancreatic duct encircling and extending to the right side of the duodenum is identified by MRCP or ERCP, the diagnosis of annular pancreas is established. But sometimes the pancreatic duct without dilatation is invisible on MRCP. Secretin-enhanced MRCP may be the best non-invasive imaging modality for evaluating ductal anatomy. The annular pancreatic duct usually communicates with the main pancreatic duct (Fig. 11) but may drain into the intrapancreatic common bile duct, the duct of Wirsung, or the duct of Santorini. At CT and magnetic resonance imaging (MRI), a ring of pancreatic tissue surrounds the descending duodenum, in continuity with the pancreatic head (Fig. 12) (10, 11). However, a visualization of a complete ring of pancreatic tissue around the duodenum is not required for a diagnosis of annular pancreas (9). Pancreatic tissue extending in a posterolateral or anterolateral direction to the second part of the duodenum or pancreatic tissue anterior and posterior to the duodenum (‘crocodile jaw’ configuration), in the presence of a gastric outlet obstruction should raise the suspicion of annular pancreas (Fig. 13) (9). It is most likely that these patients have a thin band of pancreatic tissue, not seen at CT and MRI, incorporated in the duodenal wall.

**Ectopic Pancreas**

Ectopic (heterotopic) pancreas is described as a pancreatic tissue that lacks anatomic or vascular continuity with the normal pancreas. This incidence of this condition including

**Fig. 7.** Magnetic resonance cholangiopancreatography image reveals dominant dorsal duct (DD) with santorinicele (arrow) in pancreas divisum. D = duodenum

**Fig. 8.** Magnetic resonance cholangiopancreatography image reveals long common channel (> 15 mm) (arrows) and associated Todani type 4a choledochal cysts with intrahepatic and extrahepatic components. CBD = common bile duct, MPD = main pancreatic duct

**Fig. 9.** Magnetic resonance cholangiopancreatography image shows dorsal duct (DD) crossing anterior to common bile duct (CBD) and emptying separately into minor papilla and CBD joining with ventral duct (VD) and both entering into major papilla in patient with pancreas divisum. D = duodenum
Ectopic pancreas is most commonly (70%) located at the submucosa of the gastric antrum, proximal portion of the duodenum or the jejunum. Rarely it can also be found in the ileum, colon, appendix, mesentery, gallbladder or Meckel diverticulum. The ectopic tissue usually measures 0.5-2.0 cm in its larger diameter (rarely up to 5 cm) (12). This anomaly is usually asymptomatic and occurs as an incidental finding on gastroscopy, although complications such as ulceration, bleeding, intussusception and obstruction may develop. Rarely adenocarcinoma occurring in the ectopic pancreatic tissue may be seen (13).

At barium studies, ectopic pancreas is usually seen as a smooth, broad-based submucosal lesion in the greater curvature of the gastric antrum or in the proximal duodenum. A diagnostic feature is a central niche or umbilication, representing the orifice of the rudimentary pancreatic duct, containing a small collection of barium, seen in up to 45% of cases (12, 14). There are no specific diagnostic features at CT to differentiate ectopic pancreas from other submucosal masses.
Pancreatic Agenesis and Hypoplasia of the Dorsal Pancreas

Total agenesis of the pancreas is extremely rare and is incompatible with life. Hypoplasia (partial agenesis) results from the absence of the ventral or dorsal pancreatic anlage. Dorsal pancreatic agenesis (short pancreas) may be an isolated finding but has also been reported as a part of heterotaxia syndromes (Fig. 14) (15). In complete dorsal agenesis, the anterior head, neck, body and tail of the pancreas, the duct of Santorini and the minor papilla are absent. In partial dorsal agenesis, a variable amount of pancreatic tissue is absent but a remnant of the duct of Santorini and the minor papilla are present. Patients with dorsal pancreatic agenesis have an increased risk of diabetes mellitus because most of the islet cells are located in the distal pancreas (16).

Accessory Pancreatic Lobe

The accessory pancreatic lobe, an extremely rare anomaly, is defined as an accessory lobe of pancreatic tissue originating from the main pancreatic gland and containing...
an aberrant duct (17). The accessory lobe may be short or long, with a wide or narrow connection to the main portion of the pancreas. This anomaly is usually associated with a gastric duplication cyst and the aberrant duct communicates with the main pancreatic duct and the duplication cyst (Fig. 15) (17). Recurrent acute pancreatitis is the most common (66%) clinical manifestation of this anomaly. The underlying cause of recurrent acute pancreatitis is hypothesized to be obstruction of the pancreatic duct by viscous mucus secretions, ulcer bleeding or biliary sludge (17).

Variations of Pancreatic Contours

The head and neck portions of the pancreas may have lobulated contours, especially in the lateral aspect, mimicking a pancreatic tumor, peripancreatic metastatic tumor deposit or lymphadenopathy (Fig. 16). The attenuation or signal intensity of the lobular pancreas is the same as the normal pancreatic tissue in all phases, including unenhanced, arterial, pancreatic parenchymal and portal phases. This is the key point that helps to differentiate this condition from a pancreatic neoplasm (12).

CONCLUSION

Congenital anomalies of the pancreas and pancreatic duct are not uncommonly encountered at radiologic examinations. Radiologists should be aware of these anomalies and their variable imaging features to distinguish them from other pancreatic conditions. MRCP is the imaging modality of choice for the work-up of suspected developmental anomalies of the pancreas and pancreatic duct. It may depict noninvasively the course and drainage pattern of the pancreatic duct and can easily diagnose developmental anomalies of the pancreas. Alternatively, multiplanar reconstructed MDCT images with thin slices may allow the identification of congenital anomalies of the pancreatic duct and pancreas. Recognition of developmental anomalies of the pancreas and pancreatic duct is important because these anomalies may be a surgically correctable cause of recurrent pancreatitis or the cause of gastric outlet obstruction. An awareness of these anomalies may help in surgical planning and prevention of inadvertent ductal injury.

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