Pancreaticoduodenal artery aneurysm associated with coeliac artery occlusion from an aortic intramural hematoma

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Abstract
Pancreaticoduodenal artery aneurysms are a rare type of visceral artery aneurysm, whose rupture is associated with high mortality. These aneurysms are of particular interest because local haemodynamic change caused by coeliac artery obstruction plays an important role in their development. However, the pathophysiological mechanism of coeliac artery obstruction is not completely understood. Pressure from the median arcuate ligament is most frequently reported cause. Although it is well-known that stenosis or occlusion of the visceral vessels may be caused by aortic syndrome, reports of pancreaticoduodenal artery aneurysm associated with coeliac artery occlusion due to aortic syndrome are extremely rare. Our case indicates a new aetiology for a pancreaticoduodenal artery aneurysm and demonstrates the rapid deterioration of the patient affected.

Key words: Pancreaticoduodenal artery; Coeliac artery; Visceral artery aneurysm; Aortic dissection

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Core tip: Approximately 60% of patients with pancreaticoduodenal artery aneurysms presented with rupture have an attending mortality rate of 50%. With the development of the device and techniques,
transcatheter arterial embolotherapy has decreased the mortality to as low as 0%. Therefore, early detection and treatment is necessary to improve prognosis of the case.

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INTRODUCTION

Sutton and Lawton first reported a pancreaticoduodenal artery (PDA) aneurysm associated with coeliac axis occlusion in 1973 and since then 93 such cases have been reported[1]. Obstruction of the coeliac artery leads to collateralization of the hepatic and splenic arteries through the superior mesenteric artery (SMA). Increased blood flow in the small and fragile pancreaticoduodenal arcade vessels may contribute to aneurysm development[1,2]. Here we report what is, to the best of our knowledge, the second case in the English-language literature of a PDA aneurysm associated with occlusion of the coeliac artery due to aortic syndrome.

CASE REPORT

An apparently healthy 60-year-old man developed sudden and severe upper abdominal pain and was admitted to a local hospital. He did not feel nausea or other symptoms of visceral ischemia. There was no history of any cardiovascular disorder, connective tissue disorders, chest injury, and familial history of aortic syndrome. The aortic dissections detection risk score was 1. Initial plain computed tomography (CT) performed 5 h after onset of symptoms showed a dilated descending aorta and hyperattenuated collection located eccentrically within the aortic wall (Figure 1A). However, the attending physician could not recognize it and the cause of abdominal pain remained unclear. Therefore, contrast-enhanced CT had to be performed 6 h after the onset of symptoms, which revealed crescentic and asymmetric wall thickening of the descending aorta: Contrast is not visualized within the aortic media. Intimal flap was not detected. A diagnosis of aortic intramural hematoma (IMH) extending from the distal aortic arch to the level of the coeliac artery was made (Figure 1B-D). The root of the coeliac artery was compressed by IMH (Figures 1D and 2A). The patient was transferred to our hospital 8 h after symptom onset. This patient was not administered any drug which would affect the coagulation during the period of pre-hospital to hospital care.

On arrival, his vital signs were normal and initial laboratory data were: white blood cell count 20190/mm³, D-dimer 85.9 μg/mL, and C-reactive protein 3.0 mg/dL, with other parameters, including haemoglobin, prothrombin time, activated partial thromboplastin time, and serum amylase levels within normal limits. However, the patient lost consciousness and became hypotensive (74/40 mmHg) during transfer to the CT room 10 min after arrival in the emergency room. Contrast-enhanced CT revealed an aortic IMH and a large retroperitoneal haematoma involving the head of the pancreas and duodenum (Figure 2A and B).

The patient underwent visceral angiography for localization of the site of injury and to control bleeding. The arteriogram showed complete occlusion at the root of the coeliac artery, collateralization from SMA through the pancreaticoduodenal arcade and a saccular aneurysm of the inferior PDA (Figure 2C). The patient was, therefore, diagnosed with retroperitoneal bleeding caused by a ruptured PDA aneurysm. Selective microcoil embolization proximal and distal to the aneurysm was successfully performed. A clear CT performed at our hospital revealed anterior branch aneurism of inferior PDA (Figure 2B). The patient recovered completely and has been doing well for 5 mo.

DISCUSSION

PDA aneurysms are rare and account for 2% of all the splanchnic artery aneurysms[3]. PDA aneurysms may either be true aneurysms, resulting from a primary abnormality of the vessel wall, or false aneurysms, resulting from injury and erosion of the vessel wall, most commonly after trauma or inflammation (e.g., pancreatitis). Most cases of true PDA aneurysm are reported in patients in their 50s[1,2], suggesting a different aetiology from that of atherosclerotic aneurysms reported in the elderly. In fact, coeliac axis stenosis has been identified in 60%-75% of patients diagnosed with true PDA aneurysm[4]. In contrast, the reported incidence of coeliac axis stenosis ranges from 12.5% to 24% in the general population[5].

Obstruction of the coeliac artery leads to collateralization of the hepatic and splenic arteries through the SMA. Increased blood flow in the small and fragile pancreaticoduodenal arcade vessels is likely to contribute to aneurysm development[1]. However, the pathophysiological mechanism of coeliac artery obstruction remains to be elucidated. Pressure from the median arcuate ligament (MAL) is the most frequently reported cause[5]. Although it is well known that spontaneous aortic syndrome may cause occlusion of the aortic side branch, to the best of our knowledge, only one Japanese patient with ruptured PDA aneurysm associated with acute aortic syndrome involving the coeliac axis has been reported[6]. In that report, CT at symptom onset confirmed the presence of MAL...
Figure 1 Initial computed tomography performed at local hospital before the onset of hypotension. A: Plain computed tomography (CT) demonstrating dilated descending aorta and hyperattenuated collection located eccentrically within the aortic wall (arrow); B: Contrast-enhanced CT revealed crescentic and asymmetric wall thickening of the descending aorta: Contrast is not visualized within the aortic media. Intimal flap was not detected; C: An aneurysm at the anteroinferior pancreaticoduodenal artery (arrow); D: The occluded root of the coeliac artery (arrow).

Figure 2 Subsequent contrast-enhanced computed tomography and arteriography performed at our hospital, after the onset of hypotension. A: Multidetector computed tomography (CT) performed at our hospital showing the root of the coeliac artery compressed and occluded by the false lumen more clearly than that performed at a local hospital; B: CT scan showing a large haematoma in the right prerenal space (arrow head) and the aneurysm of anteroinferior pancreaticoduodenal artery (arrow); C: Arteriogram of the superior mesenteric artery (SMA) showing blood flow reversal from the SMA to the hepatic artery through the pancreaticoduodenal arcade, due to coeliac trunk occlusion and a saccular aneurysm of the inferior pancreaticoduodenal artery (PDA) (arrow).
with acute aortic dissection. Therefore, the authors speculated that both MAL and aortic dissection played a role of causative factor to develop the aneurysm. However, our patient showed no other factor causing coeliac artery occlusion without aortic IMH. Aortic IMH are considered as a variant of aortic dissection, accounting for 10%-30% of all acute aortic syndromes. Despite the difference in the pathophysiology, the prognosis for thoracic aortic dissections and IMH is similar. Although IMH is less likely to be associated with the branch artery occlusion compared with the typical aortic dissection, there have been some reports on ischemia due to IMH3,4.

Although the duration of the coeliac artery occlusion from aortic IMH was limited in our case, we speculate that this is a causative factor contributing to PDA aneurysm development and rupture because of an interesting characteristic of PDA aneurysms. Although most visceral artery aneurysms show a correlation between the size and the frequency of rupture, this does not apply to PDA aneurysms. Because true aneurysms of the splenic and hepatic artery, accounting for 70% of all visceral artery, rarely rupture when they are < 2 cm, it is generally believed that visceral aneurysm require a lot of time to develop and rupture. However, PDA aneurysms have not shown a clear correlation between size and propensity to rupture. Brocker et al4 reported a size variance from 3 to 40 mm (mean 15.2 mm) for ruptured PDA aneurysm. In his report, at least 24 patients bled from aneurysm < 2 cm and 14 patients bled from aneurysm < 1 cm. This suggests PDA aneurysms may rupture within short time after development. Therefore, we conclude that the celiac artery obstruction due to aortic IMH played an important role in the development and rupture of PDA aneurysm in this case, even in a limited duration of time.

It is difficult to make diagnosis of PDA aneurysms before the occurrence of rupture because of the low prevalence and nonspecific symptoms. Approximately 60% of patients with PDA aneurysms presented with rupture have an attending mortality rate of 50%. With the development of the device and techniques, transcatheter arterial embolotherapy has decreased the mortality to as low as 0%. Therefore, early detection and treatment is necessary to improve prognosis of the case.

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