A rare variation of the incomplete coeliac trunk

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During the routine dissection course, we found a rare variation of the incomplete coeliac trunk in an 87-year-old Korean male cadaver. The left gastric artery, the splenic artery and the hepatomesenteric trunk arose independently from the abdominal aorta. The detailed branching pattern of this case could not be classified by classic classification systems, but belongs to type III’ of Morita’s classification. The accurate embryological knowledge on the arterial variations is important both to anatomists and to clinicians. (Folia Morphol 2016; 75, 1: 122–124)

Key words: arterial variations, coeliac trunk, primitive splanchnic branches

INTRODUCTION

The foetal vascular system consists of a pair of dorsal aortic anlage. Each anlage gives off ventral segmental branches supplying the gastrointestinal tract. The 2 dorsal aortas fuse together to form a single dorsal aorta and the pairs of ventral segmental arteries fuse in the midline by the middle of 5th week [13]. Four primitive splanchnic branches arise from the dorsal aorta in the middle region of the embryo and form the coeliac trunk and the superior mesenteric artery (SMA). These branches are connected to the ventral longitudinal anastomotic channels in early human embryos. Retention or disappearance of parts of this primitive arterial plexus could give rise to anomalous variations of the coeliac trunk and the SMA [10, 13, 17].

Numerous reports on vascular distributional patterns including their origins, numbers, and even routes have been published and classified [1–3, 7–9], since understanding the distribution of the coeliac trunk and the SMA is essential for medical practice. The existence of a hepatomesenteric trunk has been reported with the highest incidence of 3.5% [2], which is also true in Koreans as appeared in 2.96% [15]. On the other hand, the other association with a particular pattern of the coeliac trunk has been occasionally reported on the detailed branching patterns in this region. Herein, we report a rare arterial variation of the left gastric artery, the splenic artery and the hepatomesenteric trunk independently arisen from the abdominal aorta.

CASE REPORT

During a routine dissection at Jeju National University Medical School in 2014, we found a rare arterial variation of the incomplete coeliac trunk in an 87-year-old Korean male cadaver, whose cause of death was ‘cholangiocarcinoma’.

The classic coeliac trunk was not identified, but the left gastric artery, the splenic artery and the hepatomesenteric trunk independently arose from the abdominal aorta (Fig. 1). At the level of the 12th thoracic vertebral body, the left gastric artery arose from the abdominal aorta just on the left side from the median line of its anterior wall. The external diameter of this artery at its origin was 2.0 mm, and it ran laterally to reach the lesser curvature of the stomach.

At the same level just below the origin of the left gastric artery, the splenic artery arose from the ab-
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Dominal aorta on the median line of its anterior wall just on the right side of the origin of the left gastric artery. The external diameter of this artery at its origin was 5.0 mm, and it ran laterally to reach the spleen. In the course of this artery, it had a main pancreatic branch (dorsal pancreatic artery) which ran towards the posteroinferior region of the pancreas.

The hepatomesenteric trunk arose from the anterior wall of the abdominal aorta at the inferior part of the first lumbar vertebral body, which was 11.5 mm distal from the origin of the left gastric artery. The external diameter of this trunk at its origin was 6.5 mm, and it was divided into the common hepatic artery (the external diameter at its origin, 5.5 mm) and the SMA (the external diameter at its origin, 6.0 mm). The common hepatic artery ran upward and laterally, and gave off the right gastric artery (the external diameter at its origin, 2.0 mm) and the gastroduodenal artery (the external diameter at its origin, 4.8 mm) respectively.

**DISCUSSION**

Numerous resembled anatomic variations in this region were classified by previous classical studies [1, 8, 9], and clinically modified [2, 3, 7, 15, 18]. The present case can be considered to be quite similar to the previous classification, which consists of the hepatomesenteric trunk and the gastroplenic trunk. However, this case is not suitable for detailed classification and is necessary to study deep into a mechanism of embryological arterial development although this case could be included in a certain classification [18] which has flexible subcategories — no coeliac trunk.

From embryological point of view, Morita [10] classified a variety of arrangements of origin of the coeliac trunk and the SMA as 5 types and 15 forms. The Morita’s classification for the present case is type III’: the left gastric artery, the splenic artery and the hepatomesenteric trunk (Fig. 2), which has been previously reported only 1 case [14]. Similar variations with a special reference to the hepatomesenteric trunk were reported as Morita’s type II’ — the hepatomesenteric trunk and the left gastric artery [16]

![Figure 1. Photograph of this case. The left gastric artery (1), the splenic artery (2) and the hepatomesenteric trunk (3) arose from the abdominal aorta (AA) independently. And then the hepatomesenteric trunk was divided into the common hepatic artery (4) and the superior mesenteric artery (5); 6 — the dorsal pancreatic artery; 7 — the right gastric artery; 8 — the gastroduodenal artery; 9 — the right inferior phrenic artery.](image)

![Figure 2. Schematic diagram of developmental pattern based on Tandler’s embryogenic hypothesis (A, B) and type III’ of Morita’s classification (C); AA — abdominal aorta; CHA — the common hepatic artery; CT — the coeliac trunk; HMT — the hepatomesenteric trunk; LAA — the longitudinal anastomotic arteries; LGA — the left gastric artery; PVSA — the primitive ventral splanchic arteries; SA — the splenic artery; SMA — the superior mesenteric artery.](image)
and type IV’ — the gastrosplenic trunk and the hepatomesenteric trunk [4–6, 12]. Furthermore, an unclassified case of the common hepatic artery and inferior mesenteric artery arising from the SMA was also reported [11].

The limitation of this case is the portal area, which was unclear due to the previous operation for cholangiocarcinoma. The variant common hepatic artery arising from the SMA is more important to clinicians than the lienosplenic trunk or no trunk, although the course of the common hepatic artery as pre-/retro-portal vein has been suggested an important consideration for clinicians [15]. In this context, Morita’s type III’ might have been missed and taken as a variant of the gastrosplenic trunk. In addition, this rarity on Morita’s type III’ might also be caused by clinging to an old but distinct literary of Adachi [1], in which Morita’s type III’ could be regarded as a variant of type V (form 23) or type VI (form 24) in Adachi’s classification.

CONCLUSIONS

Taken together, we found a rare arterial anomaly of the left gastric artery, the splenic artery and the hepatomesenteric trunk independently arising from the abdominal aorta. The knowledge of these arterial variations throughout the embryological base is essential for the clinicians as well as anatomists.

REFERENCES