Dorsal Pancreatic Artery—a Study of Its Detailed Anatomy for Safe Pancreaticoduodenectomy

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Abstract

Early division of the dorsal pancreatic artery (DPA) or its branches to the uncinate process during pancreaticoduodenectomy (PD) in addition to early division of the gastroduodenal artery and inferior pancreaticoduodenal artery should be performed to reduce blood loss by completely avoiding venous congestion. However, the significance of early division of DPA or its branches to the uncinate process has not been reported. The aim of this study was to investigate the anatomy of DPA and its branches to the uncinate process using the currently available high-resolution dynamic computed tomography (CT) as the first step to investigate the significance of DPA in the artery-first approach during PD. Preoperative dynamic thin-slice CT data of 160 consecutive patients who underwent hepato-pancreato-biliary surgery were examined focusing on the anatomy of DPA and its branches to the uncinate process. DPA was recognized in 103 patients (64%); it originated from the celiac axis or its branches in 70 patients and from the superior mesenteric artery or its branches in 34 patients. The branches to the uncinate process were visualized in 82 patients (80% of those with DPA), with diameters of 0.5–1.5 mm in approximately 80% of the 82 patients irrespective of DPA origin. DPA branches to the uncinate process were recognized using high-resolution CT in approximately half of the patients.

Keywords Dorsal pancreatic artery · Pancreaticoduodenectomy · Uncinate process

Introduction

Blood loss reduction by avoiding venous congestion in the resected specimen during pancreaticoduodenectomy (PD) is one of the main indications of the artery-first approach [3, 6, 8, 9, 11, 18, 19, 23]. Early division of the inferior pancreaticoduodenal artery (IPDA) along with the gastroduodenal artery is considered important for avoiding venous congestion using the artery-first approach [3, 6, 8, 9, 11, 18, 19, 23].

The dorsal pancreatic artery (DPA) exists in many cases [1, 15, 16, 24, 27]. DPA ramification is complex, and there are many individual differences in DPA anatomy. DPA branches that feed the uncinate process of the pancreas exist in many cases [1, 15, 16, 24, 27]. In such cases, to completely avoid venous congestion, DPA or its branches distributed to the uncinate process should be divided before dissecting from the portal vein (PV) and superior mesenteric vein (SMV) in the artery-first approach during PD. However, no study has investigated the significance of DPA in the artery-first approach during PD.

Preoperatively, in each case, the surgeon comprehends the pancreatic vascular anatomy necessary for PD from preoperative computed tomography (CT) data. Thus, many studies have reported the CT depiction of the peripancreatic vascular system, including DPA [4, 7, 10, 14, 20, 22]. However, CT data of DPA branch distributed to the uncinate process have not been previously reported, although classical dissection studies have reported the anatomy of the uncinate process branch of DPA [16, 27]. Thus, the aim of the present study was to investigate the anatomy of DPA and its branches to the uncinate process on current high-resolution dynamic CT, as the first step to investigate the significance of DPA in the artery-first approach during PD.
Materials and Methods

We assessed preoperative, dynamic, thin-slice CT data of 160 consecutive patients who underwent hepato-pancreato-biliary surgery between January 2016 and December 2017 at our institutions. There were 100 male and 60 female patients with a median age of 70 years (36–87). The diagnoses are pancreatic cancer in 47 patients, bile duct cancer in 23, intraductal papillary mucinous neoplasm in 22, and so on.

CT scans were performed using a 320-row multidetector CT scanner (320-MDCT; Aquilion ONE; Toshiba Medical Systems, Tokyo, Japan at Dokkyo Medical University Saitama Medical Center, Japan, and Aquilion ONE; Toshiba Medical Systems, Otawara, Japan at NTT Medical Center Tokyo, Japan). A non-ionic contrast agent (350 mgI/ml, Iomeron; Bracco Eisai Co., Ltd., Tokyo, Japan) was infused at 600 mgI/kg for 30 s prior to imaging. Early and late arterial phase images were obtained at 25 and 40 s, respectively, after starting the infusion.

The existence, origin, and ramification and anatomy of DPA, especially the existence of branches to the uncinate process, along with the caliber of the arteries that feed the pancreatic head and origin and origin of the accessory middle colic artery (AMCA) were examined. All imaging data were evaluated by two authors (TT and TN) using a diagnostic workstation (Synapse EX version 1.9; FUJIFILM Medical, Tokyo, Japan) with consensus on readings.

The institutional review boards of the Dokkyo Medical University Saitama Medical Center, Japan, and the NTT Medical Center Tokyo, Japan, approved this retrospective study, and the requirement for informed consent was waived.

Results

Among the 160 patients, DPA was recognized using preoperative dynamic thin-slice CT in 103 patients (64%) (Table 1). DPA originated from the celiac axis (CA) or its branches in 70 patients (68%) and the superior mesenteric artery (SMA) or its branches in 34 patients (33%) (one patient had two DPAs originating from both CA and SMA series) (Table 1).

The most representative ramification and the anatomy of DPA originating from CA or its branches is as follows: DPA runs caudally behind the pancreatic neck or body and the proximal part of the splenic vein (SV); rarely initially has a branch to the pancreatic head; sometimes subsequently has a branch to the pancreatic body entering from the cranial side of SV; most often subsequently has branches to the pancreatic body entering from the caudal side of SV (usually connecting to the inferior pancreatic artery); in many cases, finally descends at the left posterior side of SMV and at the right anterior side of SMA; and enters into the uncinate process (Fig. 1a). The origin of DPA from CA or its branches and the frequency of the representative branches are presented in

| DPA Origin of DPA | Early head branch from the caudal side of SV (n) | Branch to the uncinate process branch (+) (n) | Caliber (mm) |
|-------------------|-----------------------------------------------|---------------------------------------------|--------------|
| CA (n = 70a, 68%) | 23 (82%)                                       | 1.8 ± 0.5                                   | 1.1 ± 0.5    |
| SA (n = 27)       | 12 (43%)                                       | 1.6 ± 0.4                                   | 1.0 ± 0.3    |
| CA (n = 28b)      | 3 (11%)                                        | 1.8 ± 0.7                                   | 1.1 ± 0.5    |
| SMA (n = 15c)     | 20 (74%)                                       | 1.6 ± 0.3                                   | 1.0 ± 0.3    |
| SMV (n = 27)      | 12 (80%)                                       | 1.4 ± 0.5                                   | 1.0 ± 0.3    |
| r-RHA (n = 8e)    | 3 (11%)                                        | 1.8 ± 0.5                                   | 1.1 ± 0.5    |
| AIPDA (n = 1)     | 0                                             | 1.8 ± 0.5                                   | 1.1 ± 0.5    |
| MCA (n = 1)       | 0                                             | 1.8 ± 0.5                                   | 1.1 ± 0.5    |

DPA dorsal pancreatic artery, SV splenic vein, CA celiac axis, CHA common hepatic artery, SA splenic artery, SMV superior mesenteric vein, r-RHA replaced right hepatic artery, AIPDA anterior inferior pancreaticoduodenal artery, MCA middle colic artery

* One case had two DPAs from both CA and SMA series
* Including three cases with DPA from the right hepatic artery that originates from CA and passes behind the portal vein into the SMV
* Including three cases with two DPAs from the SMA series (two with both arteries from SMA and one with arteries from SMA and r-RHA)
* Including two cases with DPA from the aorta and one with no CHA
* Including one case with DPA from the replaced common hepatic artery
Table 1. The branch to the uncinate process is recognized frequently irrespective of the origin of DPA.

DPA originated from SMA at 15.2 ± 9.8 mm caudally from the origin of SMA and proximally than IPDA or the trunk of IPDA and the first jejunal artery (1st JA), except in one case, in which DPA originated from the SMA between the origins of IPDA and 1st JA. DPA usually arises from the anterior or right anterior wall of SMA, anteriorly ascends right for several millimeters, and then divides into branches to the pancreatic body and uncinate process (Fig. 1b). The origin of DPA from SMA or its branches and the frequency of the representative branches are presented in Table 1. The branch to the uncinate process from DPA originating from SMA or its branches was less frequently recognized (but not significantly) than that from DPA originating from CA or its branches (23/37 vs. 59/70). The overall incidence of the branch to the uncinate process was 80% (82/103).

The prepancreatic arterial arcade (see “Discussion”) was recognized in only three (3%) among the 103 patients with DPA. Additionally, in five cases, a thin branch, which originated from the branch to the pancreatic body entering from the caudal side of SV, ran caudally from the left to right in front of SMV at the level of the caudal edge of the pancreatic neck; however, it disappeared at the right of SMV, and prepancreatic arterial arcade could not be identified.

The calibers of DPA and the branch to the uncinate process are illustrated in Tables 1 and 2. The caliber of the branch to the uncinate process ranged from 0.2 mm to approximately 2 mm. Irrespective of the origin of DPA, the mean caliber was approximately 1 mm, and the calibers were between 0.5 mm and 1.5 mm in approximately 80% of the patients. The mean diameters of the gastroduodenal artery and IPDA were 3.4 ± 0.8 mm and 1.8 ± 0.6 mm, respectively.

AMCA arises from SMA slightly more proximally compared with the 1st JA and courses to the upper left to meet the marginal artery at the splenic flexure. Conversely, the middle colic artery originates from SMA a little more distally than the 1st JA. AMCA was observed in 32 of the 160 patients (20%), and the incidence of AMCA was not different irrespective of the presence or absence of DPA (Table 3). Among 19 patients in whom AMCA was recognized along with DPA, AMCA originated from DPA in nine and from CA in six (Table 4).

Discussion

The artery-first approach has been advocated by many pancreatic surgeons for the early determination of resectability and curability of advanced pancreatic cancer before performing an irreversible step during PD [2, 3, 5, 6, 8, 9, 13, 17, 18, 21, 23, 25, 26, 28]. The key point of this technique is dissection between the pancreatic head and SMA early during PD (i.e., the early division of IPDA). In contrast to traditional PD, in which IPDA is usually divided during the last step of resection after dissection from PV and SMV, the early division of IPDA along with that of the gastroduodenal artery helps to avoid venous congestion of the resected specimen and contributes to blood loss reduction during PD.

Therefore, many pancreatic surgeons have routinely adopted the artery-first approach in patients with periampullary diseases other than advanced pancreatic cancers to reduce blood loss during PD [3, 6, 8, 9, 11, 18, 19,
Early division of IPDA along with the gastroduodenal artery has been considered important for avoiding venous congestion using the artery-first approach. However, many surgeons may experience the following situation. After dividing IPDA at its origin and dissecting the pancreatic head from SMV and PV, at the last step of resection, one or two thin arteries need to be divided when dissecting the nerve plexus of the pancreatic head. These arteries are the branches to the uncinate process from DPA. Without prior division of these arteries, completely avoiding venous congestion of the resected specimen cannot be achieved. However, no study has investigated the significance of early division of DPA or its branches to the uncinate process in the artery-first approach during PD.

Several problems need to be resolved for investigating the significance of early division of DPA in the artery-first approach during PD. The first problem that needs to be resolved is the anatomy and frequency of the uncinate process branches of DPA, which have many individual differences with regard to anatomy and ramification. The second is examining the technical feasibility of the early division of DPA or its branches to the uncinate process, according to the origin and ramification of DPA. The third is the implementation of a prospective study examining whether early division of DPA or its branches to the uncinate process helps in reducing blood loss during PD.

This study was conducted to elucidate the anatomy and frequency of the uncinate process branches of DPA detected using current high-resolution dynamic CT. The incidence of DPA has been reported to range from 64% to 100% in anatomical and radiological studies [1, 4, 7, 10, 14–16, 20, 22, 24, 27]. According to Michels [16], DPA is characterized by a course that crosses behind the proximal part of SV. However, the pancreatic magna artery, a large superior pancreatic branch of the splenic artery, may be erroneously recognized as DPA, especially in angiographical studies. This may explain the reported previously high incidence of DPA. The incidence of DPA (64%) identified in the present study corresponds to the minimum incidence previously reported.

Woodburne and Olsen [27] and Michels [16] reported in their dissection studies that DPA typically has two right branches and a left branch. This left branch corresponds to the branch entering the body in the present study. In their reports, a right branch, after producing the uncinate branch, turns across the pancreatic head to form a prepancreatic arterial arcade with a small left branch of the anterior superior pancreaticoduodenal artery. In the present study, this

| Table 2 | Distribution of the caliber of the uncinate process branch |
|---------|----------------------------------------------------------|
| Origin of DPA | Caliber of uncinate process branch |
|          | ≤ 0.5 mm | > 0.5 mm, ≤ 1.0 mm | > 1.0 mm, ≤ 1.5 mm | > 1.5 mm |
| CHA (n = 23) | 2 (9%) | 8 (35%) | 12 (52%) | 1 (4%) |
| SA (n = 22) | 3 (14%) | 12 (55%) | 6 (27%) | 1 (5%) |
| CA (n = 14) | 2 (14%) | 5 (36%) | 7 (50%) | 0 (0%) |
| SMA (n = 23) | 3 (13%) | 7 (30%) | 8 (35%) | 5 (22%) |
| Total (n = 82) | 10 (12%) | 32 (39%) | 33 (40%) | 7 (9%) |

CHA common hepatic artery, including one case with DPA from the right hepatic artery that originates early near CA and passes behind the portal vein, SA splenic artery, CA celiac axis, including one case with DPA from the left gastric artery from the aorta, SMA superior mesenteric artery, including four cases with DPA from the replaced right hepatic artery and one case with DPA from the middle colic artery

| Table 3 | Accessory middle colic artery |
|---------|-------------------------------|
| Accessory middle colic artery | (+) | (−) |
| Total | 32 (20%) | 128 (80%) |
| DPA (+) | 19 (18%) | 84 (82%) |
| DPA (−) | 13 (23%) | 44 (77%) |

DPA dorsal pancreatic artery

| Table 4 | Accessory middle colic artery (AMCA) from the dorsal pancreatic artery (DPA) |
|---------|---------------------------------------------------------------|
| DPA origin | AMCA from DPA | AMCA from SMA (not from DPA) |
| CHA (n = 28) | 3 | 2 |
| SA (n = 27) | 3 | 3 |
| CA (n = 15) | 0 | 4 |
| SMA (n = 24) | 3 | 0 |
| r-RHA (n = 8) | 0 | 1 |
| Total | 9 | 10 |

a CHA common hepatic artery, including three cases with DPA from the right hepatic artery that originates early near CA and passes behind the portal vein
b SA splenic artery
c CA celiac axis, including two cases with DPA from the left gastric artery (one from the aorta and one with no CHA)
d SMA superior mesenteric artery; 24 cases with 27 DPA

e r-RHA replaced right hepatic artery, including one case with DPA from the replaced common hepatic artery
prepancreatic arterial arcade was recognized in three cases; however, the branch forming the prepancreatic arterial arcade originated from the branch to the pancreatic body. The same anatomical thin branch was recognized in five cases additionally, but the prepancreatic arterial arcade could not be identified. This difference may be associated with the small caliber of this artery or with the limited resolution capacity of the current fine CT scans, which is affected by imaging timing. In cases with the prepancreatic arterial arcade, this branch can be easily divided in front of SMV early during PD.

Few studies have reported the frequency of individual branches of DPA [15, 16, 24, 27]. Woodburne and Olsen [27] reported that a prepancreatic arterial arcade had an incidence of 93.3% among 90% DPA occurrences. In a dissection study, Matsumura [15] showed that the uncinate branch had an incidence of 60.9% among 88.8% DPA occurrences. The 80% incidence of the branch to the uncinate process in our study is between these values.

The caliber of DPA branch to the uncinate process has never been reported. In the present study, the caliber ranged from 0.2 mm to approximately 2 mm and was between 0.5 and 1.5 mm in approximately 80% of the patients. It is known that a large caliber is associated with a strong influence on venous congestion. The significance of early division of DPA or its branches to the uncinate process will differ according to the caliber.

The feasibility of the early division of DPA or its branches to the uncinate process depends on the extent of lymph node dissection necessary along with the origin and ramification of DPA. For example, in a case in which DPA originates from SMA, the preceding division of the branch from DPA to the uncinate process with preservation of the branches to the pancreatic body is not very difficult using the supracolic anterior approach [8]. Additionally, in a case in which DPA originates from the splenic artery, the early division of DPA at its origin and preceding division of the branches from DPA to the pancreatic body are practically possible in a patient with pancreatic head cancer. It is noteworthy that venous congestion cannot be completely avoided without dividing branches from DPA to the pancreatic body before dissection from SMV and PV, even after dividing DPA at its origin. However, this technique is not practical for a patient with low-grade malignancy. To ascertain the significance of early division of DPA or its branches to the uncinate process in the artery-first approach during PD, a prospective study is required. However, the possible frequency of early division of DPA or its branches to the uncinate process according to disease variety and DPA anatomy should be examined before such a prospective study.

AMCA, which is also designated as the superior left colic artery [12], should be kept in mind as one of the anomalies concerning DPA, although it has little significance in the artery-first approach during PD.

To sum up, using the currently available high-resolution dynamic CT, DPA was identified in 64% of total patients included in the study, and branches to the uncinate process were visualized in 80% of those with DPA. In approximately 80% of the patients, the calibers of the branches to the uncinate process from DPA are between 0.5 and 1.5 mm irrespective of DPA origin.

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### Author Contribution
T Tatsuoka: Data collection, Manuscript writing.
T Noie: Project development, Data collections and management, Data analysis, Manuscript writing/editing.
T Noro: Project development, Critically revising manuscript.
M Nakata: Data collection, Manuscript writing.
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### Compliance with Ethical Standards

#### Conflicts of Interest
The authors declare that they have no conflict of interest.

#### Ethical Approval
All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.

#### Informed Consent
The requirement for informed consent was waived by the institutional review boards of Dokkyo Medical University Saitama Medical Center, Japan, and the NTT Medical Center Tokyo, Japan.

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