A significant vascular variant in oncologic pancreaticoduodenectomy: the arc of Buhler

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Abstract

Background: The arc of Buhler (AOB), a rare anastomosis connecting the superior mesenteric artery (SMA) to the celiac trunk (CA), was found in a patient suffering from an adenocarcinoma of the pancreatic head.

Case presentation: Oncologic pancreaticoduodenectomy required resection of the AOB to achieve complete tumor removal. After an uneventful clinical course in the first days, the patient suffered a severe complication. Due to ischemia of the stomach and spleen, complete resection of the stomach, spleen, and remaining pancreas had to be performed.

Conclusions: The hemodynamic impact of this arterial variant has been discussed mainly for liver perfusion, which remained intact at all times in our case. Because of the serious obstacles mentioned above, we strongly recommend that the presence of AOB be considered in preoperative diagnosis and preservation when possible. If the AOB is likely to be ligated, stenosis of the SMA or CA should be excluded and resolved before surgery.

Keywords: Arc of Buhler (AOB), Arterio-arterial anastomosis (shunt), Superior mesenteric artery (SMA), Celiac trunk (CA), Gastroduodenal artery (GDA), Pancreaticoduodenectomy

Background

Variants and anomalies of the abdominal visceral vessels are frequently found in patients [1]. However, an arterio-arterial anastomosis between the SMA and the CA, called the arc of Buhler (AOB), is present in only less than 3% of cases [2]. The first description of this variant dates from 1904, identified by Bühler and Tandler [3]. The AOB is thought to be a relic of ventral longitudinal anastomosis of the 11th and 13th intersegmental arteries during embryogenesis. In healthy, asymptomatic individuals, this collateral vessel is presumed to fulfill a significant hemodynamic role in approximately half of the cases [4]. However, in most reported cases, total or near-total stenosis of the CA is also found, making perfusion of the liver, stomach, and spleen potentially AOB-dependent [1]. Therefore, concerns have been raised about AOB in upper GI surgery. To our knowledge, we hereby report the first case involving the resection of the AOB during abdominal surgery.

Case presentation

A 66-year-old patient was admitted to the hospital because of painless jaundice and pathologic elevated cholestasis parameters. Conventional sonography revealed a malignancy-suspect tumor of the pancreatic head with dilatation of the common bile duct. Subsequently, endoscopic retrograde cholangiopancreatography was conducted with papillotomy and stenting of the common bile duct. No malignant cells were detected in the biopsies taken, although the suspected diagnosis of a malignant pancreatic head tumor was supported by endoscopic ultrasound. Contrast-enhanced computed tomography of the thorax and abdomen showed no metastases, and therefore curatively intended pancreaticoduodenectomy was evaluated.

3D reconstruction of the upper abdominal vessels from CT data was performed to plan the surgical treatment.
There was no evidence of local irresectability, but an arterio-arterial shunt was discovered originating from the dorsal part of the SMA and leading to the common hepatic artery (CHA), just after its branching from the CA (Fig. 1). The vessel joined the SMA and celiac trunk independently of the common arterial anastomosis, therefore it was considered to be the AOB. Furthermore, the celiac trunk appeared to be stenotic. After critical discussion with our radiology department, preoperative stent implantation was not performed as the stenosis of the CA was considered marginal.

A median arcuate ligament compression syndrome (Dunbar syndrome) could also have been the cause of the described stenosis. Because of the leading oncologic diagnosis, no further investigation of asymptomatic CA stenosis was performed.

The AOB showed close proximity to the tumor and possible tumor infiltration (Fig. 2). We planned pancreaticoduodenectomy with the intention of preserving the AOB if technically and oncologically feasible.

Intraoperatively, we found a locally advanced tumor with regional inflammatory changes. After mobilization of the duodenum and pancreatic head, the gastroduodenal artery (GDA) was ligated.

As the dissection progressed toward the pancreatic body, the AOB was identified cranial to the tumor.

Further caudal tumor infiltration could not be excluded, making preservation of the shunt for oncologic resection impossible. After clamping the AOB close to its termination at the CHA, there was no evidence of ischemia of the liver or stomach. This was verified by Doppler ultrasonography. The AOB was resected. The proper hepatic artery (PHA) showed a steady pulse at all times. Pancreatic anastomosis was constructed as a pancreaticogastrostomy. Single-loop reconstruction was performed for bilio-jejunal and gastro-jejunal anastomosis.

The early postoperative course was uneventful. On postoperative day 2, a planned gastroscopy showed no evidence of anastomatic insufficiency or ischemia. Having already received regular meals, the patient developed abdominal pain and elevated infection laboratory markers on postoperative day 9. An immediate CT scan of the abdomen indicated an insufficient pancreatic–gastric anastomosis. Emergency relaparotomy revealed gastric ischemia with insufficiency of the pancreatic–gastrostomy and gastrojejunostomy and necrosis of the remnant pancreas. In addition, the splenic artery (SA) showed a relatively small diameter and thrombosis. We performed gastrectomy, splenectomy, resection of the remaining pancreatic and reconstruction as esophagojejunostomy. The patient stabilized quickly with ICU-treatment. After drainage of the pleural effusion, treatment of wound dehiscence, and wound infection, the patient was discharged from the hospital on postoperative day 44.

The result of histologic examination confirmed complete resection of a 37-mm, moderately differentiated ductal adenocarcinoma of the pancreas with infiltration of the duodenum and common bile duct (TNM: pT2, pN 1 (3/21), L 1, Pn 1, G 2, R 0). There was no evidence of tumor infiltration of the resected arterial shunt.

Fig. 1 Identification of the arc of Buhler (white arrowheads, highlighted green) before surgery by a three-dimensional reconstructed computed tomography. CA celiac artery, CHA common hepatic artery, SA splenic artery, GDA gastroduodenal artery, SMA superior mesenteric artery, LGA left gastric artery, PHA proper hepatic artery, SPDA superior pancreaticoduodenal arteries, IPDA inferior pancreaticoduodenal arteries
Discussion

For the successful performance of surgical procedures in the upper gastrointestinal tract, it is important to recognize and consider vascular anomalies, especially in pancreatic surgery [5]. Although the AOB was first described in 1904, the rare occurrence of this anastomotic artery and the relatively low prevalence of pancreatic disease make an implication rare. A recent review showed that only 53 cases of the AOB have been described [1]. Considering the patient from our clinic and a recent case report, the number has increased to 55 [6].

Using PubMed for a literature review with the keywords “pancreaticoduodenectomy” and “arc of buhler”, we could identify only four cases involving pancreaticoduodenectomy and the coincidental presence of an AOB (Table 1).

In all four cases the AOB was preserved during surgery. In two cases, the vessel was identified before surgery so that a planned approach could be undertaken [5, 6]. These patients suffered from adenocarcinomas of the

Table 1  Previous reports of surgical pancreaticoduodenectomy in patients with a present arc of Buhler

| Author (year) | Condition | Surgical treatment | Postoperative course |
|--------------|-----------|--------------------|----------------------|
| Templin (2020) [6] | Adenocarcinoma, papilla vateri T3 N2 (7/13) M0 | Pancreaticoduodenectomy, preoperative identification of the AOB by digital 3D reconstruction | Mild cholangitis, no further complications |
| McCracken (2018) [8] | Intraductal papillary mucinous neoplasm, pancreatic head (suspected adenocarcinoma) | Pancreaticoduodenectomy, intraoperative AOB detection by angiography | TRANSIENT transaminits day 2, discharged day 7. Brief rehospitalization due to Klebsiella bacteremia |
| Kageyama (2016) [5] | Adenocarcinoma, ampulla of Vater T7 N0 M0 | Pancreaticoduodenectomy preoperative identification by computer tomography | Uneventful, discharged day 14 |
| Ochoa (2016) [7] | Pancreatic ductal adenocarcinoma T3 N0 M0 | Pancreaticoduodenectomy, AOB identified after surgery due to complications | Intraluminal bleeding on day 5—> IR intervention. Possible partial left hepatic lobe infarct, bacteremia, discharged day 19 |
ampulla of Vater, which had different locoregional conditions compared with the case reported here. One author reported identifying the shunt postoperatively because of bleeding on day 5 after pancreatectoduodenectomy of a ductal adenocarcinoma [7]. McCracken reported an accidental discovery of AOB during surgery [8]. The patient showed stenoses of the CA and SMA. Due to the coincidental presence of the AOB, no stent or arterial bypass procedure was required after GDA resection.

The discussion in the case reports above emphasized the importance of AOB preservation for liver perfusion. This results from the frequently associated CA stenosis, especially when no other collateral hepatic artery arises from the SMA. In contrast, our patient did not show signs of liver malperfusion at any time; however, major complications from ischemia of the stomach and spleen occurred. The vascular reconstruction of the postoperative CT scan (postoperative day 9) revealed a tremendous loss of vascularization regarding the CA (Fig. 3). Compared to the liver, under normal circumstances the stomach has a distinct blood supply consisting of more than four major arteries. The right gastroepiploic artery arises from the gastroduodenal artery (GDA) and receives regular collateral blood flow through the SMA from functional anastomoses between the pancreatoduodenal arteries. During pancreatectoduodenectomy, ligation and transection of the GDA and right gastroepiploic artery is mandatory. After dissection of the latter vessels and the AOB, our patient developed compromised blood supply to the stomach and spleen. This gastric ischemia was not evident on staged gastroscopy on postoperative day 2. Thus, we are left to speculate whether and to what extent CA stenting could have prevented ischemia of the stomach, spleen, and pancreatic remnant at postoperative day 9. Knowing the further course, retrospectively the evaluation of such an intervention would have been useful both preoperatively and early postoperatively.

**Conclusion**

In the presence of an AOB, caution should be exercised during pancreatic or hepatobiliary surgery. Preoperative angioplasty and stenting should be considered if there is evidence of CA or SMA stenosis. In circumstances requiring AOB resection, liver and gastric perfusion should be critically assessed. Surgeons should be prepared to extend the abdominal resection and possibly perform further vascular reconstruction.

**Abbreviations**

AOB: The arc of Buhler; SMA: Superior mesenteric artery; CA: Celiac trunk; GDA: Gastroduodenal artery; GI: Gastrointestinal; CT: Computer tomography; CHA: Common hepatic artery; PHA: Proper hepatic artery; LA: Splenic artery; ICU: Intensive care unit.

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**Authors’ contributions**

The authors participated in diagnostic and treatment of the patient and equally contributed to drafting the manuscript. The final manuscript was approved by all authors. All authors read and approved the final manuscript.

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All data generated or analyzed during this study are included in this published article. Further material and information on this case report are available from the corresponding author on reasonable request.

Declarations

Ethics approval and consent to participate
Ethics approval on this case report was not applicable. Standard surgical procedures and therapy were conducted independent of the academic findings. Data and observations were collected upon completion of treatment.

Consent for publication
For the publication, written informed consent was obtained from the patient about this case report.

Competing interests
The authors declare that they have no competing interests.

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