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DOI: 10.5603/FM.a2020.0052

Article type: CASE REPORTS

Submitted: 2020-03-22

Accepted: 2020-04-29

Published online: 2020-05-18

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Rare combined variations of the celiac trunk, accessory hepatic and gastric arteries with co-occurrence of double cystic arteries: a case report

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Abstract

Many variations of the celiac trunk and hepatic or gallbladder arterial supply have been reported before in many cadaveric and radiologic studies. In this case we present combined anomalies observed in dissected cadaver of a 73-years old female. The left gastric artery arises directly from the abdominal aorta and gives two branches: the right inferior phrenic artery in the proximal part and the accessory left hepatic artery in the distal part. The celiac trunk is bifurcated into the common hepatic artery and the splenic artery. The right gastric artery emerges from the left hepatic artery. The right hepatic artery gives two cystic arteries and the accessory right hepatic artery is noticed arising from the posterior superior pancreaticoduodenal artery. The deep cystic artery and the right inferior phrenic artery give hepatic branches. Also, we noticed small accessory biliary duct going to the cystic duct. This complexity of the arterial supply with anomaly of the biliary ducts have many surgical implications which will be herein discussed.

Key words: anatomical variations, accessory hepatic artery, gastric artery, double cystic arteries, right inferior phrenic artery
Introduction

The importance of the arterial supply of the liver, gallbladder and upper gastrointestinal tract is noticed in plenty of surgical procedures. J.F. Calot (1891) was the first, who emphasized the position of the cystic artery during cholecystectomy [7]. Adachi (1928) elaborated the classification of variations of the celiac trunk, which is also used currently [1] and Michels was one of the first researchers who described and classified the variability of the hepatic arterial supply [23]. According to recent reviews the most common variation in types of the incomplete celiac trunk was the hepatosplenic trunk present in 3.88% of cases with the left gastric artery originating from the abdominal aorta in 99.87% [35]. The right gastric artery most frequently goes from the proper hepatic artery – in 53% of cases [11]. Accessory or replaced hepatic arteries are observed in 30.8%. The accessory left hepatic artery which occurs in 8.2% mostly rises of the LGA (7.9%) and the accessory right hepatic artery described in 5.6% of cases with the most often origin from the superior mesenteric artery in 5.4% [17]. Double cystic arteries (or multiple cystic arteries) are observed in 8.9% [3]. The right inferior phrenic artery the most frequently originates from the abdominal aorta [32]. Our case comprises more or less frequent variations of all mentioned arteries with accessory branches to liver what makes it unique and allows to show the significance of them in clinical practice for example in transplantology, cholecystectomy or surgical oncology.

Case report

During a routine dissection for teaching purpose of the 73-year old female 10% formalin-fixed cadaver many multiple anomalies were observed. In the abdominal cavity any previous surgical interventions were not noted. At the level of Th12-L1 the LGA (4.74 mm in diameter) arises separately from the abdominal aorta (18.15 mm in diameter) curving a geniculate flexure and branches off the RIPA (2.75 mm in width) at 8.51 mm from the origin and heading up gives the second branch i.e. the aLHA (2.89 mm in diameter) after 25.15 mm in length course from the proximal branching. The small hepatic branch (1.17 mm in diameter) arises from the RIPA at 18.18 mm distal from the beginning. The aLHA gives small esophageal branch only 1.30 mm in diameter (Fig.2). The hepatosplenic trunk (5.33 mm in width) originates from the abdominal aorta 4.39 mm below the LGA and divides into the CHA and the SA. The CHA (4.67 mm in width) courses along the upper margin of the pancreas and branches into the PHA and the GDA at 27.41 mm from the beginning. The
PHA (3.68mm in diameter and 9.07 mm in length) ends up with bifurcating into the RHA and the LHA. The LHA (2.87 in width) branches to the RGA (1.79 mm diameter) at the 2.40 mm from the origin and the RGA runs down 25.94 mm to the lesser curvature crossing the CHA above (Fig.1). The RHA (3.34 mm in diameter) passes behind the CHD at the 15.46 mm from the beginning of its course and gives rise to the superficial CA (1.63 mm in width) 3.20 mm after crossing the CHD (diameter 7.07 mm), gives small branch to the liver and at the 5.24 mm from the first branching and gives the deep CA (1.48 mm in diameter) which from also arises a small hepatic branch (1.05 mm in width) (Fig. 3). The GDA at the 16.14 mm from the origin feeding branches to the right gastroepiploic artery (2.62 mm) and the anterior superior pancreaticoduodenal artery (1.95 mm diameter) and then the GDA runs down as the posterior PSPD along the head of pancreas giving the proximal branch (at the 16.37 mm from the previous branching), travels 9.89 mm and divides into the aRHA and the distal branch (1.93 mm). The aRHA (2.10 mm in diameter and 40.32 mm in length) runs parallel to the common bile duct and travels to the biliary fossa dividing into two branches before entering the liver. In the vicinity of this area the small accessory biliary duct (0.52 mm in width) going directly from V segment of the liver and at 14.58 mm from the origin merges to the cystic duct in the midway of its course (Fig. 4). Diameters of observed abnormal arteries were summarized in Table 1.

In this case, the arterial blood supply for the liver was supported by five arteries: RHA, LHA, aRHA gave off the PSPD, aLHA from the LGA, RIPA and the small branch arising from the dCA.

Discussion

The variations of the incomplete celiac trunk were repeatedly described by researches. As we mentioned, the hepatosplenic trunk was observed in 3.88% of cases with the abdominal aorta as the origin feeding of the LGA in 99.87%. More rarely the LGA arises from the SMA (0.76%) or it is absent (0.38%) [35]. The prevalence of the cases where the LGA gave off the RIPA is estimated of 2% to 4.1% [25]. Some of the researches distinguished the gastrophrenic trunk [16,22,26,28,32,37] as the common origin of the IPA and the LGA. The majority of the used classifications do not perceive it as a trunk and in most cases it is recognized in association of presence with the hepatosplenosenteric trunk [16,22,26,28] but the co-occurrence with the hepatosplenic trunk also was described [32,37].
According to the classification proposed by Whitley et al. [35], this type of trunk is not included but we came to the conclusion that it is important to mention this fact. Inferior phrenic arteries more frequently origin asymmetrically [5]. Right inferior phrenic arteries commonly arise from the abdominal aorta (49%) or the celiac trunk (41%), less frequently from the LGA or renal arteries (5.5%) [4,5,19]. Aslaner et al. noted higher frequency of the RIPA arising from the renal artery with co-occurrence of the incomplete celiac trunk [4].

The anomaly of the RGA arising from the LHA was observed by Yamagami et al. in 25.3% [36] of cases and the Eckmann et al. estimated the frequency to 15% [11].

Variations of the hepatic arteries were commonly described by many authors [9,17,23]. In the review presented by the Cirrochi et al. [9] the aLHA originating from the LGA was noticed in 3.60% (694/19284) of cases (including Michels’ and Hiatt’s findings). Jin et al. estimated the prevalence of the aLHA from the LGA as the origin to 8.2% (total cases 10211). The aRHA was observed in 5.6% of cases. Although the cases of aRHA arising from the GDA were revealed in approximately 0.04% (4 of 10211 cases which also included branches of the GDA) [17], the pancreaticoduodenal artery branched the aRHA in some of them [2]. Juszczak and Sobolewski et al. presented similar interesting case wherein also both accessory hepatic arteries were observed, aLHA arising from the LGA which arose independently from the abdominal aorta and aRHA from SMA as the feeding origin but contradistinctively to our case the PHA did not divide into two hepatic arteries. Also, they observed both inferior phrenic arteries originating from the hepatosplenic trunk (positioned opposite to the vertebral L1 level) which both branched off superior adrenal arteries [18].

Andall et al. revealed the weighted percentage of the multiple cystic arteries circa 8.9% with the highest noted number of arteries as four [3] but some researchers found this variation in 30.2% [24] and 28.3% [6] of cases. Small accessory biliary duct emerging from the liver parenchyma found in our case is classified as cystohepatic duct and the incidence of its variation (also including the cholecystohepatic duct which directly drains to the gallbladder) ranges between 0.2 and 2.3%. This variation might occur with normal CHD and CBD anatomy likewise in our case [30,31]. Frequency of observed variations were presented in Table 2.

Variations observed in our case reflect processes during fetal development. The most common variation is trifurcation of the celiac trunk (LGA, CHA and SA) separately to the SMA which is generated by interruption of anastomosis between third and fourth ventral
mesenteric root [33]. The interruption between 1. and 2. mesenteric roots with termination of anastomosis between 3. and 4. roots cause occurrence of LGA independently originating from abdominal aorta with hepatosplenic trunk. Higher prevalence of some variations of hepatic arteries also could be explained by analyzing the embryological development. Primarily the LHA arises from the LGA and the RHA from SMA and absence of the PHA which originates from the CHA after its bifurcation. If hepatic branches of the LGA or SMA with division PHA into RHA and LHA are preserved, these branches are named as accessory left hepatic artery and accessory right hepatic artery respectively. This fact clarifies high frequency of aLHA originating from the LGA and aRHA from the SMA. Grugacz et al. reported case wherein all fetal origins of hepatic arteries were preserved with presence of the middle hepatic artery feeding IV segment of the liver [14]. In our findings we observed abnormal primary right hepatic artery arising from branch of the GDA transitioned during development to the aRHA. Mahajan et al. presented similar case of preserved fetal pattern of liver’s blood supply, also they reported occurrence of esophageal branch arising from replaced LHA which emerged from LGA [21].

Doubled cystic arteries with co-occurrence of accessory hepatic arteries were reported by some authors. Loukas et al. described case wherein a small accessory CA arose from PSPD with aLHA originating from LGA which emerged independently from abdominal aorta and the hepatosplenic trunk also was observed [20]. Polgúj et al. revealed variant of GDA which gave off both accessory CA and aRHA which also gave off CA. No other anomalies of celiac trunk and hepatobiliary vasculature were observed in this case [27]. Dolenšek described occurrence of doubled CAs (larger sCA and small dCA) which both arose from aRHA. Similarly to our case both accessory hepatic arteries were present but the aRHA emerged from the SMA and the PHA was trifurcated into left, middle and right hepatic arteries with coexistence of classical trifurcation of the celiac trunk [10]. According to Andall et al. the prevalence of CA arising from abnormal RHA (replaced or accessory) is estimated to 5.58%. In this research the frequency of GDA and PSPD as origin feeding of CA was 1.94% and 0.07% respectively [3].

Vascular abnormalities described in this case are significant in surgical practice. Using the Critical View of Safety method (CVS) during laparoscopic cholecystectomy the surgeon has to precisely indentify vascular structures in the hepatobiliary triangle. In cases, wherein the origin feeding of the cystic artery is different than the RHA (LHA or GDA) or course of the cystic artery is anteriorly to the common hepatic duct, there is a high risk of ligation. Also,
in variant of multiple cystic arteries one of the CAs could be overlooked leading to uncontrolled bleeding [3]. Although the postcholecystectomy bile leaks occur in 0.2-2% of cases, the surgeon should be careful with presence of the subvesical ducts during the procedure, especially if this duct drains to the cystic duct [29,31].

Knowledge about variations of accessory or replaced hepatic arteries is crucial in planning intraarterial chemotherapy in treatment of cancers or metastases (mainly metastases of the colorectal cancer) in liver because it is essential to find appropriate artery feeding occupied lobe by neoplasm to accurately arrange hepatic arterial infusion pump (HAIP) placement and avoid infusion to arteries supplying the other organs. This procedure also include the embolisation of the extra-hepatic arteries, such as RGA, pyloric artery and GDA [8,13] and if accessory or replaced hepatic arteries arising from the embolized artery was not recognized it could cause hepatic ischemia. In procedures of chemo-embolisation and radio-embolisation it is significant to consider blood circulation in interested segment provided by the extra-hepatic arteries including inferior phrenic arteries which usually supply the I,II and VII segments. LGA and CA are also hepatic feeders, II, III segments and peri-vesicular region respectively [13]. The RIPA is mentioned as one of the main extrahepatic collateral arteries which are supplying the hepatocellular carcinoma (HCC) and usually in this case the diameter of the RIPA is larger than the diameter of the LIPA. The width of the RIPA feeding neoplasm in the liver ranged from 2 to 3.2 mm [5] and in another research the diameter greater than 2.5 mm was considered as indication of present collateral circulation [4].

Vascular anatomy of liver is also significant in transplantology, especially in living-donor liver transplantation, wherein right (adult recipient) or left lobe (pediatric recipient) is harvested because thorough analysis of liver vessels enables to preserve proper blood circulation in retained donor’s lobe and guarantees appropriate liver regeneration. In surgical practice preservation of all found anomalies of hepatic arteries is preferred [8,34] but some of them could be difficult to find during the procedure, for example the accessory left hepatic artery arising from the LGA [34]. Also, anomalies of the hepatic biliary vascularisation should be verified (especially in harvesting the right lobe) to avoid bile leakage in recipients which leads to graft rejection [8,12].

Conclusions

Anatomical variations of the arterial blood supply of upper part of the abdomen have great importance in many surgical procedures and they are strongly associated with
embryological development. Acquaintance about branching patterns of cystic arteries and hepatic arteries is significant in planning cholecystectomy or liver transplantation and in these procedures existence of accessory biliary duct should be concerned to avoid postsurgical complications. Abnormal branches to liver or gallbladder arising from gastroduodenal artery or posterior superior pancreaticoduodenal artery are rarely observed and knowledge about them facilitate better understanding about vascularisation of these organs.

**Abbreviations:** aLHA – accessory left hepatic artery, aRHA – accessory right hepatic artery, CA – cystic artery, dCA – deep cystic artery, sCA – superficial cystic artery, CBD – common biliary duct, CD – cystic duct, CHA – common hepatic artery, CHD – common hepatic duct, GDA – gastroduodenal artery, IPA – inferior phrenic artery, LGA – left gastric artery, LHA – left hepatic artery, PHA – proper hepatic artery, PSPD – posterior superior pancreaticoduodenal artery, RGA – right gastric artery, RHA – right hepatic artery, RIPA – right inferior phrenic artery, SMA – superior mesenteric artery

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Table 1. Summarized measurements of abnormal arteries in our case.

| Artery  | Diameter [mm] |
|---------|---------------|
| LGA     | 4.74          |
| RGA     | 1.79          |
| RIPA    | 2.75          |
| aLHA    | 2.89          |
| aRHA    | 2.10          |
| dCA     | 1.48          |
| sCA     | 1.63          |

Table 2. Prevalence of observed in our case variations.

| Variation                                           | Frequency (%) |
|-----------------------------------------------------|---------------|
| RIP A from LGA                                      | 2 – 4.1       |
| Hepatosplenic trunk (LGA from abdominal aorta)      | 3.87          |
| RGA from HA                                         | 15 – 25.3     |
| aLHA from LGA                                       | 3.6 – 8.2     |
| aRHA from GDA                                       | 0.04          |
| Multiple CAs                                        | 8.9           |
| Cystohepatic duct (including cholecystohepatic duct)| 0.2 – 2.3     |
Figure 1. Complex variations of the celiac trunk. LGA arises independently from the abdominal aorta and branches to the RIPA which gave off small hepatic branch and in the distal course from the LGA arises the accessory left hepatic artery. RGA originates from the LHA. aLHA – accessory left hepatic artery, GDA – gastroduodenal artery, LGA – left gastric artery, LHA – left hepatic artery, PHA – proper hepatic artery, RGA – right gastric artery, RHA – right hepatic artery, RIPA – right inferior phrenic artery, SA – splenic artery
Figure 2. Emphasized small branches to the esophagus and the liver. aLHA – accessory left hepatic artery, LGA – left gastric artery, RIPA – right inferior phrenic artery, SA – splenic artery.
Figure 3. A, B. Cystic arteries with branches to the liver arising from the dCA and RHA.
dCA – deep cystic artery, sCA – superficial cystic artery, LHA – left hepatic artery, aRHA – accessory right hepatic artery, RHA – right hepatic artery
Figure 4. Accessory RHA emerges from the posterior superior pancreaticoduodenal artery and travels parallel to the common bile duct to the gallbladder fossa, dividing into right and left branch. Small accessory biliary duct (also known as cystohepatic duct) goes directly from the liver (V segment) and reaches the cystic duct at its halfway.