The Impact of Hepatic Arterial Variations and Reconstructions on Arterial Complications in Liver Transplantation

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Abstract

Background: The purposes of the study were to determine the variations in hepatic arterial supply, to delineate the optimal methods of arterial anastomoses and reconstructions in liver transplantation and to analyse the incidence of arterial complications. Methods: The surgical anatomy of the extrahepatic arterial vascularization was investigated retrospectively in 209 donors and patients who underwent liver transplantation at Fundeni Clinical Institute (Bucharest, Romania) from January 1, 2015 to December 31, 2017. The vascular anatomy of the hepatic grafts was classified according to Michels’ description and other rare variations. Results: Anatomical variants of the classical pattern were detected in 26.3% of the livers (n = 55). The most common variant was a replaced right hepatic artery arising from the superior mesenteric artery (n = 17; 8.13%), followed by a common hepatic artery from superior mesenteric artery (n = 6; 2.87%). Arterial reconstructions were reported in 97 cases (45.5%). In recipients, used sites were intermediate: common hepatic artery (CHA) in 73.8% (n = 158), distal: proper hepatic artery (PHA) or common hepatic artery/gastro-duodenal artery bifurcation (CHA/GDA bifurcation) in 16.4% (n = 35) and proximal: coeliac trunk-splenic artery-aorta (CT–SA–A) in 9.3% (n = 20) of patients. Most common reconstructions were short graft artery (CT) on the recipient CHA (n = 33, 34.02%) and long graft artery: complex reconstruction between CT and superior mesenteric artery (SMA) - accessory right hepatic artery (RHA) from SMA on CHA (n = 12, 12.37%) and right hepatic graft artery on PHA or CHA/GDA bifurcation (n = 16, 16.49%). Conclusion: The information about the different hepatic arterial patterns, as well as establishing specific methods for arterial anastomoses and reconstructions is important in the determination of better outcomes.

Keywords: hepatic arterial anatomy, hepatic arterial variations, arterial reconstructions, liver transplantation, hepatic artery thrombosis.
INTRODUCTION

Knowledge of hepatic arterial vascularization has a significant relevance for the daily practice. The patterns of hepatic arterial supply are not constant. The usual anatomy of the hepatic arterial vascularization is a common hepatic artery arising from the celiac axis, accounting for 25 to 75% of the cases. In the variant patterns, the hepatic lobes receive arterial flow through branches coming from the superior mesenteric artery (SMA), left gastric artery (LGA) or, rarely, from other arterial trunks. Since Michels published his first report, several studies have reported not only common and rare hepatic artery variants, but also different classifications. Hepatic artery thrombosis (HAT) after liver transplantation is a severe complication often requiring urgent retransplantation. Patients with HAT may present with a fulminant clinical course or a subtle and indolent course. The time of onset of HAT has been correlated with the severity of subsequent complications, with late HAT thought to have a more benign course. However, outcomes still vary considerably.

The main purpose of this study is to accurately describe the anatomic hepatic artery in 209 donors and patients who underwent liver transplantation at Fundeni Clinical Institute (Bucharest, Romania), to delineate the optimal methods of arterial anastomoses and reconstructions in liver transplantation and to analyse the incidence of arterial complications. Indeed, we believe that surgical anatomy descriptions have a greater value for surgeons than studies based on radiologic images or autopsy findings.

MATERIALS AND METHODS

We analysed a retrospectively database of 209 patients who underwent liver transplantation at Fundeni Clinical Institute (Bucharest, Romania) from January 1, 2015 to December 31, 2017. Monitoring was carried out until April 2019 with a median follow-up of 34 months (range: 16–52 months). Liver grafts were from deceased donors (n= 187) or living donors (n= 22). Patients under the age of 18, patients with liver transplantation (LT) before the study period and retransplantation during the study period as well as patients with missing data were excluded. The main criterion considered as complication was HAT (early - occurring within 1-month post-LT or late – occurring after 1 month post-LT), defined as a lack of arterial flow determined by Doppler ultrasound and confirmed by a computed tomography scan (CT scan). Preoperative evaluation, surgical data, postoperative complications and 90-day mortality were recorded.

After a preoperative visit including a CT scan, all cases were discussed in the multidisciplinary transplantation meeting to assess the vascular anatomy of the recipient and discuss technical options. As recommended, the patency and size of the hepatic artery (HA) as well as the identification of any variation that could result in reduced flow of the HA were systematically analyzed with a radiologist.

All liver transplantsations were performed using a surgical technique under this sequence: (1) a piggyback caval anastomosis (three-vein technique), (2) an end-to-end portal anastomosis, (3) an arterial anastomosis and (4) a duct-to-duct biliary anastomosis. No porto-caval shunt was used. A sequential revascularization was done, and the graft was first perfused via the portal vein and then via the hepatic artery. In the absence of any hemorrhagic complications and/or low platelet count, all patients received prophylactic low molecular weight heparin. Acetylsalicylic acid was reserved for small arteries (diameter<5 mm) and/or complex arterial reconstructions.

Postoperative arterial patency was examined by Doppler ultrasound every day during the first week...
routinely. Subsequent examinations were carried out as indicated clinically.

The arterial reconstruction was planned before the transplantation to reduce prolonged warm ischemia time (number, size and quality of the donor and recipient arteries). In case of inadequate HA flow (small hepatic artery, intraoperative intimal dissection, mass ligation of hepatic pedicles in patients with portal cavernoma), dissection of the common hepatic artery (CHA) and then of the recipient celiac trunk (CT) was performed to determine the future site of arterial anastomosis. On a case-by-case basis, an adapted and oriented arterial anastomosis with intima-to-intima opposition was fashioned under a surgical loop with a running or separated 6-0, 7-0 or 8-0 polypropylene suture.

**STATISTICAL ANALYSIS**

Variables were compared between patients with and without arterial complications with Fisher’s exact test, Student t test or Mann-Whitney test, as appropriate. Odds Ratio (OR 95%CI) effect size was estimated by the logistic regression model and all variables with p<0.20 in univariate analysis were included in the multivariate analysis with Wald’s backward selection method. In order to account for the different lengths of follow-up in the cohort, survival curves were estimated with the Kaplan-Meier method and compared with Log-rank tests. Hazard Ratio (HR 95%CI) was estimated by the Cox regression model. The follow-up was decided as the number of months between the surgery and the last follow-up or the date of death. Statistical analyses were performed using IBM SPSS Statistics version 26.0 for Windows and IBM SPSS Statistics version 27 for MacOS. A p value < 0.05 was considered statistically significant.

**RESULTS**

**Demographic and operative data**

During the study period, 209 patients with primary liver transplantation involving whole organ graft or LDLT were included. Men were detected in 61.4% of the non HAT/HAS (hepatic artery thrombosis/ hepatic artery stenosis) group and in 85.5% of the HAT/ HAS group. The patients of the HAT/HAS group were 8 years younger than the non HAT/HAS group. All the patients presented a Child–Pugh score C. No difference was found concerning recipient age, gender, MELD (Model for End-Stage Liver Disease) score, HCC (hepatocellular carcinoma) as a cause of liver disease, erythrocyte transfusions and major complications (p>0.05). Arterial variations and arterial reconstructions were also not significantly associated with HAT/ HAS (p = 0.373 and respectively p = 0.703). The HAT/ HAS group had longer operative time than the non HAT/HAS group (p=0.007).

| Recipient characteristics | non HAT/HAS (n=202) | HAT/HAS (n=7) | p value |
|---------------------------|---------------------|--------------|---------|
| Age*                      | 50.5±11.25          | 42.0±16.5    | 0.053   |
| Gender M                  | 124 (61.4%)         | 6 (85.7%)    | 0.258   |
| HCC                       | 55 (27.1%)          | 1 (14.3%)    | 0.677   |
| Child score               | C                   | C            | -       |
| MELD score**              | 18 (6 - 35)         | 16 (10 - 21) | 0.566   |
| Operative time (hours)**  | 6.3 (4 - 13.2)      | 8.7 (5.6 - 10.3) | 0.007   |
| Blood products (U)**      | 2 (0 - 32)          | 2 (0 - 9)    | 0.783   |
| Arterial anatomy anomalies| 50 (24.8%)          | 3 (42.9%)    | 0.373   |
| Arterial reconstructions   | 90 (44.6%)          | 4 (57.1%)    | 0.703   |
| Major complications***     | 50 (24.8%)          | 4 (57.1%)    | 0.075   |
| Aortic conduit            | 5 (2.5%)            | 0 (0.0%)     | 1.000   |
| Vascular graft            | 6 (3.0%)            | 0 (0.0%)     | 1.000   |
| Aortic patch              | 7 (3.5%)            | 0 (0.0%)     | 1.000   |
| Short graft artery        | 31 (15.3%)          | 2 (28.6%)    | 0.305   |
| Long graft artery         | 10 (5.0%)           | 2 (28.6%)    | 0.054   |
| Median follow-up (months) | 35 (0 – 52)         | 23 (0 – 51)  | 0.656   |
| 90-day mortality          | 26 (12.9%)          | 2 (28.6%)    | 0.238   |

* The data are presented as medians, standard deviations. ** The data are presented as medians and ranges. *** Major complications - including grade Clavien Dindo IIIA, IIIB, IV, V.

The protective effect of the age was the same in univariate and multivariate analysis (p=0.064 and p=0.118). Major complications increased 4 times (95%CI: 0.88-19.73) the odds of HAT/HAS in univariate analysis and 3.2 times (95%CI: 0.58-17.76) in multivariate analysis, without statistical significance (p=0.073 and p=0.179). Long gaft artery increases 9.6 and 5.5 times the odds of HAT/HAS in univariate and multivariate analysis (p=0.017 and 0.072). Each operative hour significantly increases (69%) the odds of HAT/HAS (OR=1.69; 95%CI: 1.16 - 2.49; p=0.007), being also
statistically significant in the multivariate analysis (OR=1.63; 95%CI: 1.12−2.43; p=0.016).

**ARTERIAL ANATOMY AND TYPES OF RECONSTRUCTIONS**

Liver arterial anatomy was classic (Type I) in 154 patients (73.7%). We found arterial variations (donors and recipients) in 55 patients (26.3%): a replaced LHA (left hepatic artery; Type II) in 5 patients (2.39%), a replaced RHA (right hepatic artery; Type III) in 17 patients (8.13%), replaced left and right arteries (Type IV) in 3 patients (1.43%), accessory LHA from LGA (type V) in 5 patients (2.39%), an accessory RHA from SMA (type VI) in 4 patients (1.91%), accessory RHA and LHA (type VII) in 4 patients (1.91%), an accessory RHA or LHA and a replaced LHA or RHA (type VIII) in 3 patients (1.43%) and a CHA from SMA (Type IX) in 6 patients (2.87%), CHA separate origin from aorta (type X) in 1 patient (0.48%). There were also observed rare arterial anomalies (7 cases – 3.35%): artery for segment IV from RHA, 2 LHA from proper HA, artery for segments V-VI from GDA, CA-SMA common trunk (celiac-mesenteric trunk), early bifurcation of the HA, LHA from CA + RHA from aorta, RHA from CA.

Arterial reconstructions were reported in 97 cases (45.5%). In recipients, used sites were intermediate (CHA) in 73.8% (n = 158), distal (PHA or CHA/GDA bifurcation) in 16.4% (n = 35) and proximal (CT–SA–A) in 9.3% (n = 20) of patients. The most common reconstructions were short graft artery (CT) on the recipient CHA (n = 33, 34.02%) and long graft artery (complex reconstruction between TC and SMA - accessory RHA from SMA) on CHA (n = 12, 12.37%) and right hepatic graft artery on PHA or CHA/GDA bifurcation (n = 16, 16.49%). A donor iliac artery interposition graft was used in 6 cases (8.8%), when adequate inflow from the native hepatic artery cannot be ensured, as in very small vessels or high-grade celiac axis stenosis. Recipient aorta was used in 5 cases (7.1%). A Carrel aortic patch was performed in 7 cases (10.3%).

Arterial anatomy variations are detailed in Table 4.

**Table 2.** Uni and multivariate comparison of the HAT/HAS and non-HAT/HAS groups

| Recipient characteristics | Univariate | Multivariate |
|---------------------------|------------|--------------|
|                           | OR (CI95%) | p value      |
| Age                       | 0.95 (0.89 - 1.00) | 0.064 | 0.95 (0.89 - 1.01) | 0.118 |
| Long graft artery         | 9.58 (1.46 - 50.4) | 0.017 | 5.48 (0.86-34.98) | 0.072 |
| Operative time (hours)    | 1.69 (1.16 - 2.49) | 0.007 | 1.63 (1.10 - 2.43) | 0.016 |
| Major complications       | 4.05 (0.88 - 19.73) | 0.073 | 3.22 (0.58 - 17.76) | 0.179 |

Prominent liver transplant indications (table 3) were: (1) hepatocellular carcinoma in 56 patients (26.79 %), (2) viral cirrhosis B+D in 48 patients (22.96 %), (3) viral cirrhosis C in 29 patients (13.87 %), (4) alcoholic liver disease in 28 patients (13.39 %).

**Table 3.** Transplant indications for the HAT/HAS and nonHAT/HAS groups

| Indications                     | non HAT/HAS group (n=202) | HAT/HAS group (n=7) |
|---------------------------------|---------------------------|--------------------|
| Hepatocellular carcinoma        | 55                        | 1                  |
| Viral cirrhosis B+D             | 46                        | 2                  |
| Viral cirrhosis C               | 28                        | 1                  |
| Alcoholic liver disease         | 28                        | 0                  |
| Acute liver failure             | 6                         | 0                  |
| Budd-Chiari syndrome           | 6                         | 0                  |
| Primary sclerosing cholangitis  | 4                         | 0                  |
| Primary biliary cirrhosis       | 2                         | 1                  |
| Other indications               | 27                        | 2                  |
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Table 4. Arterial anatomy variations

| Description | Type (Michels and Hiatt) | Number of cases | %  |
|-------------|--------------------------|-----------------|----|
| standard anatomy | I | 154 | 73.7 |
| replaced LHA | II | 5 | 2.39 |
| replaced RHA | III | 17 | 8.13 |
| replaced RHA and LHA | IV | 3 | 1.43 |
| accessory LHA from LGA | V | 5 | 2.39 |
| accessory RHA from SMA | VI | 4 | 1.91 |
| accessory RHA and LHA | VII | 4 | 1.91 |
| accessory RHA and LHA and replaced LHA or RHA | VIII | 3 | 1.43 |
| CHA replaced to SMA | IX | 6 | 2.87 |
| CHA separate origin from aorta | X | 1 | 0.48 |

Rare anomalies:
- artery for segment IV from RHA
- 2 LHA from proper HA
- artery for segments V-VI from GDA
- CA-SMA common trunk (celiac-mesenteric trunk)
- early bifurcation of the HA
- LHA from CA + RHA from aorta
- RHA from CA

Total cases 209

Prominent arterial reconstructions are described in Table 5.

Table 5. Prominent arterial reconstructions

| Prominent arterial reconstructions | Cases | %  |
|-----------------------------------|-------|----|
| Short graft artery (CT) – recipient CHA | 33 | 34.02 |
| Long graft artery (complex reconstruction between TC and SMA; accessory RHA from SMA) – recipient CHA | 12 | 12.37 |
| Right hepatic graft artery – PHA or CHA/GDA bifurcation | 16 | 16.49 |
| Donor iliac artery interposition graft | 6 | 8.8 |
| Recipient aorta | 5 | 7.1 |
| Carrel aortic patch | 7 | 10.3 |

INCIDENCE AND PRESENTATION OF ARTERIAL COMPLICATIONS

Early HAT was diagnosed in 3 patients (1.43%) of the 209 patients with LT with a median delay of 8 days after LT (range: 5–15). Late HAT occurred in 2 patients (0.95%) with a median delay of 80 days (range: 40 – 120). There were described 2 cases of hepatic artery anastomotic stenosis of the 209 patients with LT. Presentation of the arterial complications included elevated transaminases, biliary complications (biliary leakage), fever, abnormal results on Doppler images (abnormal RI), sepsis and graft dysfunction or failure.

MANAGEMENT OF HAT, OUTCOMES AND SURVIVAL ANALYSIS

Of the 7 patients with arterial complications, all patients received medical treatment (acetylsalicylic therapy), 4 patients underwent endovascular treatment with Actilyse and thrombectomy and 2 patients underwent retransplantation (at POD 10 and 46).

The mortality of patients with arterial complications was 28.6% (n =2), while the mortality of patients without arterial complications was 17.3% (n =35). Mean survival time in the HAT/HAS group was 43.9 months versus 36.9 months in the nonHAT/HAS group, without statistical significance (p=0.365). The risk of mortality during the follow-up is increased in the HAT/HAS group: HR=1.91 (95%IC: 0.46 - 7.93), compared with the nonHAT/HAS group.
The mortality of patients without arterial anomalies was 19.2% (n =30), while the mortality of patients with arterial anomalies was 13.2% (n =7). Mean survival time in the arterial anomalies group was 45.5 months versus 43.1 in the standard anatomy group, with no significant difference (p=0.366). Arterial anomalies were a protector factor for mortality during the follow-up (HR: 0.69; 95%CI:0.30 -1.56).

The mortality of patients with standard anastomoses was 22.6% (n =26), while the mortality of patients with arterial reconstructions was 11.7% (n =11). Mean survival time in the arterial reconstructions group was 46.2 months, versus 41.8 in the standard anastomoses group (p=0.069). Arterial reconstructions were a protector factor for mortality during the follow-up (HR: 0.53; 95%CI: 0.26 - 1.07).

DISCUSSION

Our main findings can be summarized as follows: the prevalence of anatomical variants was 26.3%, a percentage similar to that reported in other series \(^4\); most of the variants described fit into Michels' classification; the most common anatomical variants in our series were a replaced right hepatic artery (type III) in 17 patients (8.13%), in agreement with previous studies and a CHA from SMA (type IX) in 6 patients (2.87%). The findings of the present study highlight the fact, already observed in previous studies, that the extrahepatic arterial distribution is variable and that different anatomic variants can occur in a high percentage of cases. These arterial patterns are relevant in the procurement of donor livers and also in the planning and performance of all types of liver transplantations \(^7\).

We considered it important to identify all the hepatic arteries and to classify them as "accessory" or "replaced". Indeed, replaced arteries must be always preserved; in contrast, accessory arteries do not necessarily need to be reconstructed if there is adequate backflow after the anastomosis of the other branch or if arterial flow in all the segments is demonstrated by intraoperative Doppler ultrasonography \(^6\).

The analysis of these data shows the predominance of type I (no arterial variations, a single hepatic artery, which originates in the celiac trunk): 154 cases (73.7%). 10 cases (4.78%) had left hepatic artery (replaced or accessory) as left gastric branch, and 21 (10.04%) had a right hepatic artery (replaced or accessory) originating from the superior mesenteric artery. 6 cases (2.87%) presented a single hepatic artery originating from the superior mesenteric. 7 cases were not described in the Hiatt classification. The most common reconstruction was CT on the recipient CHA (n = 33, 34.02%) and long graft artery (complex reconstruction between TC and SMA - accessory RHA from SMA) on CHA (n = 12, 12.37%) and right hepatic graft artery on PHA or CHA/GDA bifurcation (n = 16, 16.49%). A donor ili-ac artery interposition graft was used in 6 cases (8.8%), when adequate inflow from the native hepatic artery cannot be ensured, as in very small vessels or high-grade celiac axis stenosis. Recipient aorta was used in 5 cases (7.1%). A Carrel aortic patch was performed in 7 cases (10.3%).

Arterial variations and arterial reconstructions were not significantly associated with HAT/HAS (p = 0.373
and respectively p = 0.703). The HAT/HAS group had longer operative time than the nonHAT/HAS group (p=0.007). Major complications increased 4 times (95%CI: 0.88-19.73) the odds of HAT/HAS in univariate analysis and 3.2 times (95%CI: 0.58-17.76) in multivariate analysis, without statistical significance (p=0.073 and p=0.179). Long graft artery increases 9.6 times and 5.5 times the odds of HAT/HAS in univariate and multivariate analysis (p=0.017 and 0.072). Each operative hour significantly increases (69%) the odds of HAT/HAS (OR=1.69; 95%CI: 1.16 - 2.49; p=0.007), being also statistically significant in the multivariate analysis (OR=1.63; 95%CI: 1.12-2.43; p=0.016). The mortality of patients with arterial complications was 28.6% (n =2), while the mortality of patients without arterial complications was 17.3% (n =35). The risk of mortality during the follow-up is increased in the HAT/HAS group: HR=1.91 (95%IC: 0.46 - 7.93), compared with the non HAT/HAS group.

While transplantation is the most effective treatment for acute and chronic end-stage liver disease, it is also a procedure with significant morbidity and mortality, particularly in the early postoperative period. This crucial time can encompass several complications, the most significant being primary graft nonfunction and hepatic artery thrombosis. Hepatic artery thrombosis is the second main cause of liver graft failure. The consensus for early HAT definition consists of an arterial thrombosis detected during the first month after liver transplantation. HAT is associated with markedly increased morbidity, being the leading cause of graft loss (53%) and mortality (33%) during the immediate postoperative period. However, improvements in postoperative care have resulted in a marked reduction of its incidence.

Early hepatic artery thrombosis was diagnosed in 3 patients (1.43%) of the 209 patients with LT, with a median delay of 8 days after LT (range: 5–15). Late hepatic artery thrombosis occurred in 2 patients (0.95%) with a median delay of 80 days (range: 40 – 120). There were described 2 cases of hepatic artery anastomotic stenosis. Of the 7 patients with arterial complications, all patients received medical treatment (acetylsalicylic therapy), 4 patients underwent endovascular treatment with Actilyse and thrombectomy and 2 patients underwent retransplantation (at POD 10 and 46).

The clinical expression depends on the timing of the onset of HAT as well as on the existence of collateral arteries. The natural history of early HAT is biliary tract necrosis followed by uncontrolled sepsis in the immunosuppressed population and even death. Late HAT can be asymptomatic, but in most cases the clinical expressions are biliary tract complications such as necrosis and abscess, as well as liver graft ischemia. The key point in vascular complications is early diagnosis, which allows immediate treatment to avoid graft loss. Doppler ultrasonography is the gold standard for screening protocols, because it detects the absence of hepatic artery flow, even in its intrahepatic branches. The screening protocols or early HAT between different centers are highly variable with respect to frequency. In cases of an early suspicion of HAT, diagnostic plus therapeutic arteriography should be performed.

The clinical features depend on the type of complications (thrombosis, stenosis, pseudoaneurysm), timing of occurrence (early or late presentation after liver transplantation), and promptness of the diagnosis. Treatment options include surgical revascularization, percutaneous thrombolysis, percutaneous angioplasty or retransplantation.

Artery complications represent one of the main causes of graft failure and death after liver transplantation. In addition to parenchymal damage, decreased hepatic artery perfusion can lead to biliary complications such as necrosis, the formation of bilomas, bile leaks, and the development of non-anastomotic structures. Sonographic demonstration of these findings or of parenchymal abscesses should raise the suspicion of ischemia. A strict surveillance program with doppler ultrasonography can allow early detection of artery complications and prompt therapy. The most common causes of thrombosis are the vessel folding in on itself because of excessive length, organ rejection, elderly donors, slow flow, stenosis, artery reconstruction, and previous liver transplantation. Most stenoses originate at or within a few centimeters of the arterial anastomosis. Several surgical causes in the development of the hepatic artery stenoses have been proposed, such as trauma to the hepatic artery (clamp injury and/or intimal dissection), hepatic artery kinking, differences in vessel caliber requiring oblique anastomoses, and extrinsic compression. Other causes include microvascular preservation injury, disruption of the vasa vorum of the allograft hepatic artery, and acute cellular rejection.

The key point in vascular complications is early diagnosis, which allows treatment to avoid graft loss.
Doppler ultrasonography is the gold standard for screening protocols. The use of US color Doppler examination allows the early diagnosis of hepatic arterial complications after liver transplantation. Tardus-parvus waveforms indicated severe impairment of hepatic arterial perfusion from either thrombosis or severe stenosis. The presence of these indirect signs enhanced the accuracy of color Doppler diagnosis, and detection should prompt therapy.

In many instances, HAT occurs soon after surgery for technical reasons, but HAT can also occur several months after transplantation due to factors such as graft rejection or sepsis. Z. Stewart et al showed that donor age is associated with late graft loss induced by HAT. As described by J. Bekker et al, the overall 1-year graft survival rate after liver transplantation for early HAT is approximately 50%, while other vascular complications have graft survival rates up to 86%.

The incidence of hepatic arterial occlusion ranges from 4% to 9% requiring a second liver transplant in over 50% of patients. Christian E. Oberkofler et al. found that primary arterial patency after LT is predominantly determined by the type of vascular reconstruction rather than patient or disease characteristics.

The ideal arterial reconstruction is often described as a short and non-redundant anastomosis fashioned between the recipient and donor hepatic arteries. A. Herrero et al. found in multivariate analysis that the use of a long graft artery, whatever the recipient anastomosis site, was an independent risk factor of early HAT (OR 3.2; 95% CI 1.2–9; p = 0.02). Arterial reconstruction using a long graft artery on distal recipient arterial site (PHA or CHA/GDA bifurcation) was associated with an increased rate of early HAT compared with reconstruction on other sites (p = 0.02). No difference was found between recipient sites of implantation when using a short graft artery.

The favourable outcome of LHAT compared to early arterial thrombosis is allowed by the development of arterial collaterals, that can arise from superior mesenteric artery, splenic artery, inferior phrenic artery, left gastric artery and arteries from the omentum, providing arterial blood supply via the hilar plate. This could explain the persistence of resistive index in patients with no more patent hepatic artery.

Marudanayagam R. et al. concluded that the most common indication for retransplantation was hepatic artery thrombosis (31.6%). Nearly two-thirds of the retransplantations were performed within 6 months of the primary transplantation. The 1-, 3-, 5- and 10-year patient survival rates following first retransplantation were 66%, 61%, 57% and 47%, respectively. Donor age of <55 years and a MELD score of <23 were associated with better outcome following retransplantation (being independent risk factors predicting outcome following regrafts). Univariate analysis was performed initially to identify factors predicting survival. Early retransplantation was associated with worse outcome compared with late retransplantation (p = 0.015).

The timing of retransplantation has been shown to play a significant role in both patient and graft survival. Chen et al. demonstrated that patients in whom the interval to retransplantation was 8–30 days displayed lower survival rates compared with those who underwent later retransplantations.

Compliance with ethics requirements: The authors declare no conflict of interest regarding this article. The authors declare that all the procedures and experiments of this study respect the ethical standards in the Helsinki Declaration of 1975, as revised in 2008(5), as well as the national law. Informed consent was obtained from all the patients included in the study.
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