Chapter 14
Liver Transplantation

Michael Sean Bleszynski and Peter T. W. Kim

Introduction

For a critically ill cirrhotic patient, liver transplant is the only treatment that can provide a chance at long-term survival. In patients who meet the criteria for transplant listing, the current allocation system is designed to direct the next available donor liver to the sickest patient on the list to reduce wait-list mortality. Liver allocation was originally based on overall wait times, the Child-Turcotte-Pugh (CTP) score, and ABO blood type compatibility [1]. However, this allocation scheme had limitations in that longer waiting times on the transplant list did not correspond with increased patient mortality and the CTP score did not adequately represent the general transplant population [1]. The CTP score is based on three laboratory values (prothrombin time, bilirubin, albumin) and two subjective clinical variables (ascites and encephalopathy). Despite the CTP scoring model initially being utilized as part of an organ allocation system, it has never been validated for estimating survival in patients with chronic liver disease [1]. The CTP score is rather reflective of complications of portal hypertension, and its lack of objectivity limited its application to transplant organ allocation [2].

The model for end-stage liver disease (MELD) score was initially developed to determine risk of mortality for the transjugular intrahepatic portosystemic shunt (TIPS) procedure within a 3-month period [3]. The MELD score has subsequently been validated as a severity of liver disease scoring system and predictive mortality tool independent of etiology or occurrence of portal hypertensive complications [1, 4].
Baseline MELD scores have been shown to be significantly associated with wait-list mortality [4]. Since its approval in 2002, the MELD score helps determine liver allocation for patients awaiting transplantation by providing 3-month predictive mortality [5] and has become the most commonly utilized liver organ allocation tool worldwide. Sicker patients are represented by a higher MELD score and therefore are assigned a higher priority on transplant waiting lists. The main advantage of the MELD score is that it is objective in that it is based on three laboratory values (serum INR, bilirubin, and creatinine). It is not a perfect system in that it doesn’t always reflect the urgency in patients with relatively low physiologic MELD score but who have clear indications for liver transplantation (e.g., hepatocellular carcinoma, hepatopulmonary syndrome, metabolic disorders) [6]. These patients are usually granted MELD exception points that would help them to be competitive for transplants depending on their region of residence. More recently, the MELD-Na has been introduced to provide a more accurate assessment of wait-list mortality and to take into account the complications of portal hypertension [7].

Despite the advancements within transplantation over the last 20 years, several challenges remain, as organ shortages persist and patients remain on wait-lists for extended periods of time. Due to the current allocation system based on MELD, transplant programs are often offering liver transplants for patients with high MELD scores. This raises new challenges and questions in today’s practice. This chapter aims to outline current evidence for transplantation of patients with high MELD scores, discuss transplant futility, address simultaneous multi-organ transplantation, discuss surgical techniques for complications of cirrhosis at the time of transplantation, discuss postoperative management, and outline the role of living-related transplantation in today’s environment.

Liver Transplantation in High MELD Patients

The MELD score has been validated as a scoring tool to prioritize patients on liver transplantation (LT) waiting lists by predicting 3-month mortality risk based on a scale from low scores of 6 to high scores capped at 40, with 83–87% accuracy [1]. Wait-list mortality is directly proportional to the MELD score, where a MELD score of <9 is associated with an approximate mortality of 2% and a MELD score ≥ 40 is associated with a wait-list mortality of 71% [1]. In general, for the patients with MELD scores ≤15, the risks of LT likely outweigh the benefit. In low MELD patients, the risk of mortality from LT is greater than remaining on the transplant wait-list [8]. These patients are therefore allocated to the bottom of the list and, depending on the program, often not listed until their MELD score increases. Application of the MELD score has reduced the number of patients awaiting LT and lowered peri-transplant mortality [6]. The MELD score has not been able to improve the shortage of available organs. Organ distribution is based on medical urgency, rather than expected posttransplant outcomes. Patients with MELD scores >35 are typically admitted to the ICU, potentially on dialysis, receive hemodynamic or
respiratory support [9], and are potential candidates for urgent LT. In such situations, it may seem that sick patients would not benefit from operative intervention. However, based on a 5-year time frame, the higher the MELD score, the greater the benefit of LT [10]. Survival benefit posttransplantation is seen in MELD scores >40 because this population has the greatest risk of mortality while awaiting LT [11].

Patients with MELD scores >40 were previously thought to be “too sick” to undergo LT. It was believed that organ allocation to this higher risk population was futile and not beneficial for individual patient outcomes or for appropriate resource utilization. Currently, the pretransplant MELD score has not been able to reliably predict posttransplant outcomes [12, 13]. As patients linger on waiting lists, MELD scores continue to increase. It is common to see patients with MELD scores >40 awaiting LT. Low-MELD-score patients may also spend a prolonged amount of time on wait-lists, deteriorate, and become part of the sickest quartile of individuals awaiting LT. Interest lies in assessing which critically ill patients with high MELD scores derive the most benefit from LT. In patients with MELD scores >40, are there additional factors not captured by the MELD score that can predict successful or futile transplantation outcomes? In order to reduce wait times, in 2013, the United States adopted the Share 35 policy, which mandated that there would be an increase in regional sharing of organs to patients with MELD scores ≥35 [14].

A Canadian retrospective review assessed the outcomes of 198 critically ill ICU cirrhotic patients undergoing LT with a median MELD score of 34 on ICU admission [15]. The 90-day and 3-year survival were 84 and 62.5%, respectively, despite the fact that 88% of patients received vasopressors, 56% received renal replacement therapy, and 87% were mechanically ventilated prior to transplantation [15]. The same study found that patients >60 years of age had a significantly higher 90-day mortality (27% vs 13%) [15]. A multivariate analysis of 8070 transplant patients aged ≥60 identified that recipient albumin levels <2.5 mg/dL, serum creatine ≥1.6 mg/dL, hospitalization at the time of organ offer, ventilator dependence, presence of diabetes, and recent hepatitis C virus (HCV) positivity were independent predictors of poor patient survival [16]. In this study, the strongest prognostic factor was a recipient and donor age combination equal to or greater than 120 years [16]. Asrani et al. [17] retrospectively reviewed non-HCV cirrhotic LT recipients and identified that patients who had a survival of <50% at 5 years were above 60 years old with median MELD scores of 40. These patients also had multiple medical comorbidities and were on life support at the time of LT. Age > 60 in patients with elevated MELD scores has consistently been shown to be associated with worse posttransplantation survival compared to those patients with MELD scores >40 and age < 60.

Patients with MELD scores >40 have increased wait-list mortality compared to patients with lower MELD scores [1]; however, an elevated MELD score is no longer a contraindication to LT [18]. Studies have shown contradictory results for high MELD score patients and postoperative mortality. Retrospective analysis has demonstrated that cirrhotic patients with MELD scores ≥40 do benefit from LT and have similar 5-year cumulative survival posttransplantation compared to patients with MELD scores <40. However, these patients confer a higher burden of health-care
In 2014, the University of California, Los Angeles, group showed similar findings, in which patients with MELD score > 40 LT was deemed beneficial with a 5-year patient survival rate above 50% [20]. The same group also identified that a subgroup of patients with MELD scores >40 did not benefit from transplantation. Patients with MELD scores >40 who had septic shock, cardiac risk factors, and other significant comorbidities, were found to have a predicted futility of LT of >75% [20]. Prospective analysis has confirmed the association of elevated healthcare costs with MELD scores ≥28 is due to longer hospital and ICU admissions, despite no differences seen in postoperative survival or complications when compared to MELD scores <28 [18].

Panchal et al. [21] retrospectively reviewed a nationwide transplant database and found that the overall mortality was statistically higher in patients with MELD scores ≥40, compared to patients with a MELD score < 30 (30 versus 26%). Despite the significant difference in mortality, the MELD >40 group had a lower mortality rate than initially predicted, which was thought to be secondary to younger age of recipients, lower prevalence of diabetes, portal vein thrombosis (PVT), HCV, Epstein-Barr virus, TIPS, or prior upper abdominal surgery [21]. This group utilized greater hospital resources (longer pretransplant hospitalization, ICU admission, required mechanical ventilation, and longer hospital length of stay) [21]. Within the same study, MELD patients with a score of >40 and recipient age > 60, BMI > 30, pretransplant hospitalization, or use of extended criteria donors predicted LT futility. The risk of mortality increased by 95%, and graft failure was 60% higher when compared to patients with a MELD <30. Despite the significantly increased risk, there is a perceived benefit to transplanting such sick patients because they have expected survival of >50% at 5 years (64% graft and 69% patient survival) [21]. In recipients with satisfactory graft function, MELD scores >30 are significantly associated with prolonged ICU stay (defined as ≥3 days) which is associated with poor patient and graft survival at 3, 12, and 60 months [22]. However, good LT outcomes can be seen in patients with MELD scores ≥40, where overall 1-, 3-, 5-, and 8-year survival of 89, 79, 75, and 69% can be seen when futile deaths are excluded [20].

There is a definite subgroup of patients in this high-risk category in whom LT becomes futile despite optimal management. Michard et al. performed a single-center retrospective review and identified that in patients awaiting LT with MELD scores >40, those admitted to the ICU had elevated lactate (>5 mmol/L) or developed acute respiratory distress syndrome (ARDS) and had a poor 3-year survival rate of 29% [23]. In this subpopulation, LT is clearly not beneficial. A comparison of several studies with high MELD scores and associated variables predicting poor patient survival is summarized in Table 14.1.

There are several challenges of offering transplants to patients with high MELD scores. Selecting the most appropriate donor organ for the most appropriate recipient in order to provide the best postoperative survival can be challenging. Single-center experience has demonstrated that high-risk donor organs transplanted in low MELD patients has resulted in lower recipient transplant survival [26, 27]. Furthermore, the quality of the donor organ has not impacted recipient survival in recipients with MELD scores >30 [28]. The complexity of organ allocation systems
### Table 14.1 High MELD score and variables associated with poor survival

| Study                  | Year, study type | Total number of patients | MELD | Overall patient survival | Recipient factors associated with poor survival |
|------------------------|------------------|--------------------------|------|--------------------------|-------------------------------------------------|
| Nekrasov et al. [24]   | 2017 Retrospective single center | 207 | ≥40 | 86% at 1 year 79% at 3 years 73% at 5 years | DM RRT prior to transplant Pretransplant PVT |
| Nekrasov et al. [25]   | 2016 Retrospective UNOS database | 5002 | ≥40 | 80% at 1 year 72% at 3 years 67% at 5 years 53% at 10 years | Age > 60 Hospitalization time Previous liver transplant Previous abdominal surgery Ventilator dependence HCV DM |
| Karvellas et al. [15]  | 2013 Retrospective multicenter | 198 | 34 (median) | 84% at 90 days 74% at 1 year 62.5% at 3 years | Age > 60 |
| Aloia et al. [16]      | 2010 Retrospective UNOS/OPTN database | 8070 (92% of patients between age 60 and 69) | MELD score available for 40% of cohort | MELD ≥23 75% at 1 year 72% at 3 years MELD 16–23 83% at 1 year 77% at 3 years MELD <16 87% at 1 year 77% at 3 years | Albumin <2.5 mg/dL Hospitalization Ventilator dependence DM + HCV Cr ≥ 1.6 mg/dL Combined recipient/donor age of ≥120 years |
| Asrani et al. [17]     | 2018 Retrospective SRTR/OPTN database | 31,829 | 23 (median) | 79.1% at 5 years Survival ≤50% at 5 years for; age > 60, median MELD 40, and on life support | Ventilator support Age > 60 HD Cr ≥ 1.5 mg/dL without HD DM |
| Petrowsky et al. [20]  | 2014, Single-center retrospective | 169 | 42.2 (mean) | 72% at 1 year 64% at 3 years 60% at 5 years 56% at 8 years | "Cardiac risk Age-adjusted CCI ≥ 6 Life support treatment Pretransplant septic shock" |

(continued)
Table 14.1 (continued)

| Study | Year | Database | MELD ≥40 | MELD 30–39 | MELD <30 | Age > 60 | BMI > 30 | ICU or ventilation | Multiple comorbidities | Obese or extended criteria donors |
|-------|------|----------|----------|------------|----------|---------|----------|-------------------|-----------------------|-------------------------------|
| Panchal HJ et al. [21] | 2015 | Retrospective UNOS database | 33,398 | 2610 patients | 5984 patients | 24804 patients | | | | |

Legend: SRTR (Scientific Registry of Transplant Recipients), OPTN (Organ Procurement and Transplantation Network), HD (hemodialysis), Cr (Creatinine), HCV (hepatitis C virus), CCI (Charlson comorbidity index), RRT (renal replacement therapy), PVT (portal vein thrombosis)

- Recipient factors associated with graft failure, rather than poor survival
- Cardiac risk defined as severe valvular disease, coronary artery disease with 70% stenosis or previous revascularization, history of myocardial infarction, history of ventricular/atrial arrhythmias, increased pretransplant troponin, new wall motion abnormality on echocardiography
- Factors associated with poor survival were analyzed in a subpopulation of patients with MELD ≥40, in order to assess predictors of futility

in individual countries can further complicate organ allocation. In 2013, the Share 35 policy was implemented in order to enhance distribution of organs in the United States for patients with MELD scores ≥35 [29]. Since its implementation, there has been a 36% reduction of organ offers accepted for patients with MELD scores ≥35, while there was no change in organ acceptance for MELD scores <35 [29]. The most common reasons for declining an organ offer were “patient transplanted, transplant in progress, or other offer being considered,” indicating that programs had several offers to choose from and were selectively choosing donors that were deemed to be more optimal [29].

As high-MELD-score patients continue to be transplanted, ongoing study is required to assess how a multidisciplinary approach with surgeons, hepatologists, and anesthesiologists can continue to enhance perioperative care in order to improve short and long outcomes. It is imperative to establish a consensus of independently successful and futile predictors of transplant outcomes in patients with MELD scores ≥35, in order to optimize outcomes in high-risk patients and prevent futile LT.

Liver Transplantation and Futility

Medical futility can divided into four major types: physiological, imminent demise, lethal condition, and qualitative [30]. When LT was in its surgical infancy prior to becoming recognized as a life-altering treatment that should be offered to patients with end-stage liver disease (ESLD), the procedure was associated with physiologic futility. Physiologic futility is defined as a proposed treatment that cannot lead to its intended physiologic effect [31] such as the case if a patient with ESLD undergoes LT and the patient does not survive the operation.
Imminent demise futility is closely associated to physiologic futility. In imminent futility, a performed action may have prolonged an individual’s life, however, only for the very short term. For example, a patient with ESLD undergoes LT, and a few days or weeks later pass without being discharged from hospital. The intent of the transplant was to extend the patient’s life by years, and the result was below this expectation. In this situation there is a subjective perceived benefit; the patient’s life was prolonged; however, the patient may or may not have believed that the short extension of life was of benefit to them. Lethal condition futility is an extension of imminent demise where the expectation is that a patient will pass away in the near term regardless of receiving or undergoing an intervention; however, the short extension on life is deemed appropriate. For example, biliary stent placement in patients with advanced incurable biliary tree tumors does not reduce mortality but provides symptom reduction thus enhancing remaining quality of life. A controversial definition of futility is qualitative futility, because it requires the scientific assessment of the probability of success for a given treatment [32].

Quantitative futility is defined by Schneiderman et al. [33], “where a treatment should be considered futile is if it has been useless in the last 100 cases, only preserves permanent unconsciousness, or fails to end total dependence on intensive medical care.” Qualitative futility addresses the end result of the intervention performed and whether the functional outcome is acceptable or not [32]. How do we as a society universally agree on what is considered acceptable? Within today’s society there are diversely held cultural and religious beliefs on what defines an acceptable quality of life outcome after an intervention in the setting of potential imminent death. A consensus definition in such a setting would be a milestone achievement. Qualitative futility encompasses the current and future ethical ambiguity surrounding transplantation of very sick, physiologically deranged patients. In today’s environment, the ethical questions and dilemmas are typically no longer dominated by the technical aspects of “can it be done?” but have transitioned to “should it be done?” Performing a highly complicated anastomosis, transplanting a patient with a MELD score > 40 with adverse prognostic indicators, or re-transplanting a patient several times, is no longer technically impossible. The ability to withdraw from aggressive medical treatment in the setting of limitless options should propagate reflection on what we consider optimal versus futile care.

How can we identify what is currently considered futile but will no longer be considered futile in the next decade of LT? Identification and stratification of patients with MELD scores >40 with associated poor predictors of outcome is necessary in order to establish a consensus of specific conditions that independently provide significant postoperative challenges that may be insurmountable to the patient. In such situations, the focus should be on the application of qualitative futility: enhancing remaining quality of life, reducing hospital resource utilization, and preserving organs that might be of a more long-term benefit to other recipients. A consensus on how we define poor outcomes in situations of ESLD and imminent death is required. How do we determine what an acceptable survival rate is, and should it be based on being better than 50%, the flip of a coin? Should there be an objective evaluation, assessing success on a minimum 5-year survival and predetermined cost? We cannot
solely focus on what is best for an individual patient. Consideration must also be
given to what is best for the next patient awaiting LT. Unfortunately, resource utiliza-
tion and medical costs are also a mandatory part of the conversation.

Many scoring systems predicting post-LT outcomes are available, and specific
definitions of futility have been created by several groups. Petrowsky et al. [20] and
Panchal et al. [21] defined futility as a 90-day mortality or in-hospital mortality in
patients with MELD scores ≥40. Petrowsky et al. [20] also identified that in patients
with MELD scores ≥40 who underwent LT, futility was significantly associated
with greater pretransplant morbidity, higher cardiac risk, age-adjusted Charlson
comorbidity index of ≥6, life support treatment, and pretransplant septic shock. In
this population, cardiac and septic causes of death were significantly higher com-
pared to patients without futility-associated risk factors and MELD scores ≥40.
Based on their observed findings, Petrowsky and his group state that despite high
medical acuity, patients with high MELD >40 without associated futile risk factors
have successful long-term survival, and therefore such patients should be trans-
planted. Asrani et al. [17], on the other hand, defined futility as any adult recipient
with a >50% mortality at 5 years posttransplant. Rana et al. [34] state that LT in any
patient with MELD score > 40 is likely futile because the predicted posttransplant
mortality is greater than any wait-list mortality as predicted by the MELD score.
However, based on previously discussed data, there are subsets of patients with
MELD scores >40 that have good posttransplant outcomes, and a general policy of
no LT for MELD scores >40 would not be appropriate.

Despite multiple proposed definitions for transplant futility, there are no global
consensus criteria that clearly define transplant futility or provide a consensus on
LT futility-associated criteria. No guidelines currently propose delisting patients
deemed futile for transplantation from wait-lists. Delisting may provide a benefit
by optimizing remaining quality of life, rather than proceeding with LT despite
poor expected outcomes. For example, should a patient with a MELD score > 40,
age > 60 with extensive cardiac risk factors undergoing dialysis, be delisted in
order to optimize organ reallocation to another individual? Would family consent
be required? What body of governance would make such a decision, and would this
be considered too paternalistic of an approach? In North America, institutions
review these unfortunate patient situations on a case-by-case basis. Multidisciplinary
conferences, where decisions regarding high-risk cases are reviewed, play an
important role in assessing not only the recipient but also the potential donor. The
Baylor College of Medicine established the Houston City-Wide Task Force on
Medical Futility, where a committee was created to preserve and protect patient
rights while establishing a fair procedural process for potentially futile clinical situ-
ations [30].

With the limited supply of organs, objective evaluation of a patient’s transplant
candidacy should also take place and assessment if optimal allocation of organs is
indeed to those critically ill patients at the top of the transplant list. Establishing a
clear set of defined criteria that warrants a patient from being delisted from a trans-
plant waiting list may help optimize organ allocation and globally improve out-
comes. Linecker et al. [35] provide general definitions of futility and propose the
concept of “potentially inappropriate” LT by risk profiling a patient’s clinical situation and probability of not surviving the early posttransplant recovery phase. If a predictive post-mortality score could be validated to accurately prognosticate post-transplant mortality risk and incorporate donor characteristics, enhanced allocation and minimization of futile transplants could occur.

Preoperative Preparation of a Sick Patient for Liver Transplantation

**Hepatorenal Syndrome**

Please refer to Chaps. 2 and 5 on this topic.

**Porto-pulmonary Hypertension**

Porto-pulmonary hypertension (POPH) is a disease where secondary pulmonary hypertension develops in the setting of portal hypertension with or without cirrhosis [36]. POPH occurs in 2–10% of all patients with cirrhosis, with approximately 1% of all patients with POPH demonstrating severe symptomatic disease [37]. The diagnosis of POPH is based on right heart catheterization findings and requires a mean pulmonary artery pressure of ≥25 at rest, an elevated pulmonary vascular resistance >240 dyne s/cm$^{-5}$, and a normal pulmonary capillary wedge pressure (PCWP) <15 mmHg [38]. Classification of mild, moderate, and severe disease is based on mean pulmonary artery pressures of >25 to <35, ≥35 to <45, and ≥45, respectively [39].

Untreated POPH is considered to be a relative contraindication for LT, and mean pulmonary pressures >35 is an absolute contraindication to proceed with LT. After reperfusion of transplanted liver, the increased venous return will exert the volume and pressure to the right heart against high pulmonary resistance resulting in right heart failure and likely death. All the potential liver transplant patients are screened with transthoracic echocardiogram where right ventricular systolic pressure (RVSP) is estimated based on the tricuspid jets. If the RVSP is found to be elevated, these patients undergo further testing with a right heart catheterization. It is important to distinguish between primary pulmonary hypertension and volume overload which can commonly occur in patients with cirrhosis. In centers that use Swann-Ganz catheters routinely in LT, this simple measurement can identify undiagnosed pulmonary hypertension prior to starting the operation, allowing the transplant team to abort the case if the pulmonary pressure is found to be too high (>35 mm Hg).

Pharmaceutical vasodilators such as prostacyclin analogues, phosphodiesterase inhibitors, and endothelin receptor antagonists lower mean pulmonary artery pressures and allow for clinical stability evidenced by improved pulmonary hemodynamics [38,
Patients with mean pulmonary artery pressure ≤ 35 and peripheral vascular resistance <400 dynes/sec/cm$^{-5}$ can be considered transplant candidates and can receive an exception MELD score of 22 points [44]. If patients do not meet transplant criteria, they can be medically treated to a mean pulmonary artery pressure < 35 mmHg and peripheral vascular resistance <400 dynes/sec/cm$^{-5}$; then MELD exception points can be provided and increased by 10% every 3 months if there is continued hemodynamic improvements [45]. POPH patients who are transplant eligible also have significant mortality potential. It has been shown that wait-list mortality or removal from the wait-list secondary to clinical decompensation is 23.2% with a median wait-list time of 344 days [46]. Age, initial MELD score, and pulmonary vascular resistance are independent risk factors for wait-list mortality [46]. Patients with the lowest wait-list mortality are those with MELD score ≤ 12 and initial pulmonary vascular resistance of ≤450 dynes/s/cm$^{-5}$ [46].

Data from the Scientific Registry of Transplant Recipients between 2002 and 2010 was retrospectively reviewed by Salgia et al., and they identified 78 out of 34,318 patients who underwent cadaveric transplantation for POHP with MELD exception points [38]. The unadjusted 1- and 3-year patient survival for recipients with POPH was 85 and 81%, while graft survival was 82 and 78% respectively. After adjusting for donor and recipient factors, POPH recipients have a significantly higher adjusted risk of death and graft failure within the first posttransplant year compared to non-POPH transplants [38]. DuBrock et al. have reported an unadjusted 1-year posttransplant mortality rate of 14% similar to Salgia et al. [46]. Rajaram et al. performed a 10-year retrospective review between 2005 and 2015 with the objective to compare posttransplant outcomes of patients diagnosed with POPH and pulmonary venous hypertension versus patients without pulmonary hypertension [47]. The authors identified 28 patients with POPH, 13 of which underwent LT with an average MELD score of 21 [47]. One patient passed away intraoperatively; 30-day survival was 92.3%, and 1-year survival was 69.2% compared to a 1-year survival of 100% in the non-pulmonary hypertension group [47]. A recent systematic review demonstrated a 1-year posttransplant mortality rate of 26% for POPH compared to 12.7% in non-POPH patients [48]. A retrospective national cohort study of 110 POPH patients in the United Kingdom identified no difference in survival between cirrhotic and non-cirrhotic patients, and the overall survival rate at 1, 2, 3, and 5 years was 85, 73, 60, and 35% [49].

Renal Failure and Liver Transplantation

Please refer to Chap. 5 on this specific topic.

An alternate treatment strategy for liver transplant candidates with renal insufficiency is to proceed with LT and assess for the development of postoperative renal
insufficiency [51]. In high MELD patients undergoing SLKT, there is a high risk of renal allograft failure. As such, it has been suggested that liver-alone transplantation should be performed with assessment at 3 months posttransplant for potential prioritization for kidney allocation [53]. Fong et al. reported that renal allograft and patient survival were significantly lower in patients undergoing SLKT compared to isolated kidney transplantation [54].

The potential benefits of SLKT has been an ongoing debate, as there is no high-quality evidence demonstrating which patients benefit most from SLKT. A cited benefit of SLKT is immune protection of the renal allograft with lower rates of acute and chronic rejection compared to sequential kidney transplant [55]. The potential drawback of SLKT is that liver recipients receive a donor kidney when their native kidney might in fact recover, resulting in a lost organ for a patient waiting for kidney-only transplantation [53]. The ultimate goal in selecting patients for SLKT is to identify which ESLD transplant candidates will develop or have irreversible kidney damage at the time of transplantation and therefore will ultimately benefit from a single operation. The difficulty lies in that there is no reliable method to identify which liver transplant candidates with concurrent kidney injury will recover renal function or eventually require a renal transplant post LT [52]. Currently, there is no universal policy for SLKT. In 2015, Puri and Eason summarized the evolution of recommendations and guidelines for SLKT outlined below [56] [Table 14.2].

Combined Liver and Thoracic Transplantation

Combined liver thoracic transplantation is a rare phenomenon. From 1995 to 2016, there have been 17 single-center published reports [58]. Combined heart and liver transplant (CHLT) is only performed at a few select high-volume centers. From 1988 to 2015, there have been 192 CHLTs performed in the United States [59]. The rate of CHLTs being performed in the United States is rapidly increasing. A retrospective review of the UNOS database between 1987 and 2010 identified 97 reported cases of CHLTs [60]. The two most common primary cardiac diagnoses were amyloidosis (26.8%) and idiopathic dilated cardiomyopathy (14.4%), while the two most common primary liver diagnoses were amyloidosis (27.8%) and cardiac cirrhosis (17.5%) [60]. Other common indications for CHLT are for patients with heart and liver failure secondary to hemochromatosis and familial hypercholesterolemia and for patients with ESLD who have severe heart disease and are unfit for liver-alone transplantation [61]. Beal et al. summarized the following number of CHLTs performed at high-volume centers within the United States: Mayo clinic (n = 33), Hospital of the University of Pennsylvania (n = 31), University of Pittsburgh Medical Center (n = 14), University of Chicago Medical Center (n = 13), Methodist Hospital (n = 13), and Cedars-Sinai Medical Center (n = 9), with the remaining centers performing ≤7 CHLT each [59].

Cannon et al. reported that liver graft survival in 97 CHLT was 83.4, 72.8, and 71% at 1, 5, and 10 years, while cardiac graft survival was 83.5%, 73.2, and 71.5%,
| Study | Recommendations for SLKT |
|-------|--------------------------|
| Nadim et al. [50] 2012 | 1. Candidates with persistent AKI ≥ 4 weeks with one of the following:  
  (a) Stage 3 AKI as defined by modified RIFLE, i.e., a threefold increase in serum creatinine (Scr) from baseline, Scr ≥ 4.0 mg/dL with an acute increase of ≥0.5 mg/dL or on renal replacement therapy  
  (b) Glomerular filtration rate (GFR) ≤ 35 mL/min (MDRD-6 equation) or GFR ≤ 25 ml/min (iothalamate clearance)  
  2. Candidates with CKD, as defined by the National Kidney Foundation for 3 months with one of the following:  
  (a) eGFR ≤ 40 ml/min (MDRD-6 equation) or GFR ≤ 30 ml/min (iothalamate clearance)  
  (b) Proteinuria ≥ 2 g a day  
  (c) Kidney biopsy showing ≥30% global glomerulosclerosis or ≥30% interstitial fibrosis  
  (d) Metabolic disease |
| OPTN Kidney Transplantation Committee and the Liver and Intestinal Organ Transplantation Committee (OPTN Policy 3.5.10) | (a) CKD requiring dialysis with documentation of the CMS form 2728  
  (b) CKD (GFR ≤ 30 ml/min) by MDRD-6 or iothalamate measurement and proteinuria >3 g/day  
  (c) Sustained AKI requiring dialysis for 6 weeks or more (defined as dialysis at least twice per week for 6 consecutive weeks)  
  (d) Sustained AKI (≤ 25 ml/min) for 6 weeks or more by MDRD6 or direct measurement not requiring dialysis  
  (e) Sustained AKI: Patients may also qualify for SLK listing with a combination of time in categories (c) and (d) above for a total of 6 weeks  
  (f) Metabolic disease |
| Eason et al. [51] 2008 | (a) Patients with ESRD with cirrhosis and symptomatic portal hypertension or hepatic vein wedge pressure gradient ≥10 mmHg  
  (b) Patients with CKD with GFR ≤ 30 ml/min  
  (c) Patients with AKI/HRS with Scr ≥ 2 mg/dL and dialysis ≥8 weeks  
  (d) Patients with evidence of CKD and kidney biopsy demonstrating ≥30% glomerulosclerosis or 30% fibrosis  
  Other criteria recommend are the presence of comorbidities such as diabetes, hypertension, age > 65 years, other preexisting renal disease along with proteinuria, renal size, and duration of elevated serum creatinine |
| Davis et al. [57] 2007 | (a) Patients with CKD with a measured creatinine clearance (or preferentially an iothalamate clearance) of ≤30 ml/min  
  (b) Patients with AKI and/or HRS on dialysis for ≥6 weeks  
  (c) Patients with prolonged AKI with kidney biopsy showing fixed renal damage  
  (d) SLK not recommended in patients with AKI not requiring dialysis |
respectively [60]. An interesting observation was that patients who received CHLT had lower rates of acute rejection compared to patients undergoing isolated heart transplantation [60]. A retrospective study from Mayo Clinic demonstrated that the incidence of T-cell-mediated rejection was 31.8% in CHLT recipients compared to 84.8% in isolated heart transplant recipients with similar overall incidence of antibody-mediated rejection [62]. Cannon et al. note that the average MELD score at the time of CHLT was 13.8; however, the wait-list mortality for these patients would have been higher compared to patients with isolated hepatic failure with similar MELD scores [60]. Between January 1997 and February 2004, there were 110 patients wait-listed for CHLT within the United States; 33 patients (30%) underwent CHLT, 30 patients (27%) died, 11 patients (10%) were still wait-listed, and 34 patients received single-organ, sequential organ transplant or were awaiting transplant of the second organ [59]. A large single-center case series from the University of Bologna reported on 14 patients with combined heart and liver transplant failure where 13 patients underwent CHLT and 1 underwent combined heart-liver-kidney transplantation. The 1-month, 1-year, and 5-year survival rates were 93, 93, and 82%, respectively, while graft free rejection at 1, 5, and 10 years for the heart was 100, 91, 36, and 100% and 91 and 86% for the liver [61].

Patients with end-stage pulmonary disease and ESLD who are not expected to survive with only a single-organ transplant can be considered for combined lung and liver transplantation (CLLT).

Isolated lung transplantation should be considered if there is a >50% risk of mortality secondarily to the primary lung disease within 2 years if a lung transplant is not performed, >80% chance of survival at 90 days after lung transplantation, and a >80% chance of 5-year post transplant survival with adequate graft function [63]. In addition to the lung transplant indications mentioned, if there is biopsy proven cirrhosis with a portal pressure gradient >10 mmHg, a CLLT can be considered [63]. Contraindications to CLLT include albumin <2.0 g/dL, INR > 1.8, presence of severe ascites, or encephalopathy [63].

Similar to CHLT, CLLT is rarely performed, and experience is limited to single-center or multicenter case reports. Double lung transplant is most often performed during CLLT instead of single lung transplant. The most common indication for CLLT is cystic fibrosis with pulmonary and liver involvement. Other indications include POPH with ESLD, hepatopulmonary fibrosis, alpha-1 antitrypsin deficiency with advanced lung and liver involvement, and sarcoidosis [64]. As with CHLT, there is a postulated immunological benefit for combined transplant, where LT is immune protective [65, 66]. There are no standardized recommendations available for CLLT, and candidacy is evaluated at each center with a multidisciplinary board committee review [64]. Potential CLLT candidates need to be placed on individual organ wait-lists. Prior to 2005, the United States and the Euro transplant region donated lungs based on patient waiting time [67]. In May 2005, the lung allocation score (LAS) was introduced, which is comprised of several patient clinical and laboratory parameters, and in the United States the LAS has replaced waiting time for determining priority of donor lungs [67]. Other European countries have followed suit over the years.
Patients undergoing CLLT derive a significant survival benefit from CLLT; however, there is a higher risk of wait-list mortality compared to single-organ transplantation [64, 68]. Survival rates are improving for CLLT. In 2008, Grannas et al. reported the largest published single-center cohort of CLLT with 1- and 5-year mortality rates of 69 and 49% [69]. Retrospective review of 14 consecutive patients who underwent simultaneous liver and thoracic transplantation included 10 patients who underwent CLLT [58]. In seven CLLT patients, the lung was transplanted prior to the liver, and three patients underwent a liver first principle while the lungs were perfused ex vivo [58]. One hundred percent of the CLLT patients were alive at 1 and 5 years with 10% suffering acute liver rejection, 40% acute lung rejection, and 10% chronic liver/lung rejection [58]. One of the largest single-center American series included 8 patients who underwent CLLT with reported patient and graft survival of 87.5, 75, and 71% at 30 days, 90 days, and 1 year [70].

CLLT can be performed with a liver first, then lung transplant approach or alternatively with a lung-first approach. Theoretical advantages of the liver first principle include reduced complications of hepatic reperfusion, potentially reduced need for blood products, reduced incidence of donor pulmonary edema, and reduced incidence of biliary strictures [58]. Advances are continuing to evolve for CLLT in critically ill patients. Extracorporeal membrane oxygenation with central cannulation has successfully been implemented after lung transplantation and prior to orthotopic LT in order to manage extensive pulmonary reperfusion edema and right heart insufficiency [71].

Intraoperative Preparation of a Critically Ill Recipient for Liver Transplant

Historically, adult orthotopic LT has been associated with massive hemorrhage with median red blood cell (RBC) transfusion rates of 28.5 units per case [72]. With improved surgical technique, intraoperative anesthetic management, transfusion medicine, and improved understanding of coagulation abnormalities [73] associated with cirrhosis, intraoperative transfusion rates have been steadily decreasing over the last 20 years [74]. Patients with low MELD scores can undergo transplantation with 0.3 units of packed RBCs without plasma, platelet, or cryoprecipitate transfusion [75], while increased INR and presence of ascites have been independently correlated with increased intraoperative blood product utilization [76, 77].

With reduced blood product transfusions, survival posttransplantation has improved [78–80]. In fact, transfusion of one or more units of plasma has been shown to have a 5.1 increased mortality risk compared to no plasma received [81]. A retrospective analysis of 286 transplant recipients found that the strongest predictor of overall survival was the number of blood transfusions after a mean follow-up of 32 months [77]. In order to identify which transplant recipients are at an increased risk of requiring intraoperative blood products, Mccluskey et al. developed a risk index score for massive blood transfusion and identified 7 preoperative variables
including age > 40 years, hemoglobin ≤10 g/dL, INR 1.2–1.99 and >2, platelet count ≤70 × 10^9/L, creatinine >110 umol/L (females) and >120 umol/L (for males), and repeat LT [82].

Normal hemostasis requires a balance between the coagulation and the fibrinolytic systems. One of the pathophysiologic complications of end-stage cirrhosis is the reduced ability or inability of the liver to synthesize new or clear activated coagulation factors [83]. During technically challenging cases, surgical bleeding can be magnified by the inability of the recipient liver to produce coagulation factors and platelets for necessary clot formation. A majority of cirrhotic patients will exhibit some form of thrombocytopenia, which is secondary to increased platelet activation, consumption, and splenic sequestration of platelets associated with portal hypertension [84]. Although total number of platelets are reduced, it has been shown that in the remaining platelets, there is increased activity secondary to increased levels of von Willebrand factor and decreased levels of ADAMTS 13 [85]. All the coagulation factors are synthesized by the liver, the only exception being factor 8. In cirrhotic patients the levels of vitamin K-dependent factors fall by 25–70% [86].

Cirrhosis induced thrombocytopenia in conjunction with prolonged prothrombin time (PT), and activated partial thromboplastin time (aPPT) was previously thought to be indicative of an increased bleeding risk [85, 86]. However, cirrhotic patients have a “rebalanced” homeostasis of anticoagulant and procoagulant cascades [85]. Furthermore, the etiology of cirrhosis can impact the balance between coagulopathy and thrombosis [83]. In the critically ill cirrhotic recipient prior to LT, superimposed infections, renal injury, endotoxins, and imbalances of coagulation factors [87] contribute to the coagulopathy seen intraoperatively. Understanding the coagulopathic profile of severely cirrhotic patients and the impact of the phases of LT is important in order to anticipate intraoperative challenges.

The initial abdominal incision made is based on surgeon preference. Commonly utilized incisions for opening the abdomen include a bilateral subcostal incision with upper midline laparotomy (Mercedes incision) or an upper midline laparotomy with a right lateral extension (Cheney incision). Table-mounted Thompson, Omni, or Bookwalter retractors are used to help facilitate intra-abdominal exposure, and choice of retractor is typically also dependent on surgeon preference. When the abdomen is opened, it is important to be cognizant of patients with ascites. Quick removal of large-volume ascites upon entering the abdomen can potentially result in a rapid shift of recipient hemodynamics.

The general steps of LT are divided into pre-anhepatic, anhepatic, and neohepatic/reperfusion phases. The pre-anhepatic phase refers to recipient hepatectomy and is completed once the vascular inflow/outflow has been controlled and clamped. Once vascular inflow and outflow have been clamped, the anhepatic phase begins, and the recipient liver is removed. The anhepatic phase continues with implantation of the new donor liver and subsequent IVC and portal vein anastomosis. The neohepatic/reperfusion phase begins with unclamping of the venous inflow and outflow, perfusion of the donor liver, and venous return to the heart. Subsequently the hepatic arterial and biliary anastomoses are performed, and the neohepatic phase is complete. Recipient warm ischemia time generally refers to the time that the recipient
liver has been explanted to the time that the donor liver has been implanted and flow through the donor graft has been established.

During the pre-anhepatic phase, the recipient liver is completely mobilized by taking down the falciform, triangular, and coronary ligaments of the liver. Once the liver has been mobilized, portal dissection is performed in order to identify and isolate the common bile duct, right/left and common hepatic arteries, and the portal vein. Dissection of the gastrohepatic ligament provides access to the portal structures. The common bile duct, right and left hepatic arteries, and portal vein are subsequently ligated. The common bile duct should be resected just distal to the cystic duct. The gastroduodenal artery should be identified; however, it does not routinely need to be ligated.

In severely cirrhotic patients with portal hypertension, the pre-anhepatic phase is usually associated with the greatest amount of bleeding. The surgeon may encounter several potentially large portosystemic collaterals in the setting of the previously described hyperdynamic circulation [88], complicating mobilization, and dissection of the recipient liver. Adhesions secondary to prior upper abdominal surgery can further complicate the hepatectomy phase [89], and previous abdominal surgery has been found to be an independent risk factor for blood transfusion requirements [90]. Reduced availability of coagulation factors and platelets inhibits the liver’s normal ability to deal with surgical bleeding.

During the recipient hepatectomy measurement and prophylactic treatment of abnormal laboratory bleeding time (BT), PT, INR, and aPTT have been common practice in order to help control anticipated surgical bleeding. However, as early as 1997, it was identified that aggressively correcting laboratory coagulation abnormalities prior to the anhepatic phase of transplantation is not required and that over-resuscitation during the pre-anhepatic phase may lead to extensive blood loss [91]. Prophylactic administration of FFP and RBCs contributes to blood loss by increasing splanchnic pressure in an already hyperdynamic circulatory state. Infusion of additional volume will eventually circulate back to the heart during the neohepatic phase [92]. As such, the utility of prophylactic treatment of abnormal laboratory values in cirrhotic patients has been questioned [84, 87, 93]. An evolving trend is the minimization of blood product transfusions during LT.

In general, there are two anhepatic techniques of LT: caval interposition and caval sparing (i.e., piggyback technique). The classic caval interposition technique begins with a retrohepatic caval dissection with cross-clamping of both the suprahepatic and infrahepatic inferior vena cava (IVC). This is followed by removal of the recipient liver, interposition and anastomosis of the donor IVC, and liver graft to the recipient suprahepatic and infrahepatic IVC. Suprahepatic and infrahepatic IVC reconstruction is performed with 3-0 or 4-0 Prolene sutures in a running fashion. Prior to completion of the infrahepatic caval anastomosis, the donor portal vein is flushed with a preservation solution in order to rid the liver of accumulated toxins that may contribute to reperfusion syndrome. Once flushing is complete, the donor portal vein is reconstructed to the recipient portal vein in an end-to-end fashion with 6-0 Prolene sutures. Typically, a shorter portal vein reconstruction is preferred over
a longer donor/recipient portal vein reconstruction with the hope of reducing kinking or development of postoperative portal vein thrombosis.

Re-establishment of blood flow with unclamping of the IVC and portal anastomosis completes the anhepatic phase, and reperfusion of the liver begins. There are alternative flushing techniques described and are based on surgeon preference. Historically, venovenous bypass was used in conjunction with classic caval reconstruction. The purpose was to provide venous return when the usual caval venous return to the heart is interrupted [94]. Nowadays, it would be commonplace to perform caval interposition technique without venovenous bypass.

Alternate caval reconstruction techniques such as the piggyback [95] or side-to-side [96] caval anastomosis only require partial occlusion of the recipient suprahepatic IVC. The recipient liver is mobilized off of the recipient IVC, while the IVC is left intact. Care must be taken while dissecting the liver off the IVC as retrohepatic veins may easily tear, cause further bleeding, and potentially damage the IVC.

In the piggyback technique, the donor hepatic vein can be anastomosed to two or three recipient hepatic veins. If only two hepatic veins are used, then the right hepatic vein is ligated. The piggyback technique with partial IVC occlusion provides a theoretical advantage of maintaining venous blood flow from the infrahepatic IVC to the heart. Maintaining cardiac preload is believed to stabilize hemodynamic stability and therefore avoids large intraoperative fluid infusions and potential need for vasopressors. Additional suggested advantages of partial caval occlusion include shorter anhepatic phase and possible decreased incidence of renal injury [97]. Moreno-Gonzalez et al. retrospectively identified that the piggyback technique was associated with longer operative times but also with less intraoperative hemodynamic instability, RBC transfusions, pressors, and fluid administration [98]. Graft outflow obstruction and increased incidence of bleeding from the caval anastomosis are recognized potential complications of the piggyback technique [97]. Caval obstruction associated with the piggyback technique is thought to be secondary to a large donor graft causing compression or an inadequate graft size that can result in twisting of the caval anastomosis, ultimately leading to hepatic venous outflow obstruction [99].

The transition from the anhepatic to neohepatic phase is critically important as there is no functioning liver during the anhepatic phase. No clotting factors are produced, and the concentration of tissue plasminogen activator increases, which contributes to fibrinolysis [100]. The accumulation of citrate leads to increased binding of ionized calcium, and calcium is an important cofactor for proper hemostasis [101]. Pooled systemic blood below the IVC clamp becomes cold and hyperkalemic as lactic acid, toxic metabolites, cytokines, and free radicals accumulate and cannot be removed [102]. When the IVC and portal vein clamps are removed, circulation is restored, and the donor liver receives the systemic blood and forwards it toward the recipient heart while the portal vein provides a fresh inflow of blood.

At this critical time, reperfusion syndrome can induce recipient hemodynamic instability as the pooled systemic blood is returned to the heart. Hilmi et al. classified postreperfusion syndrome (PRS) as mild or severe [102]. Mild PRS occurs when the decrease in blood pressure and or heart rate is <30% of the anhepatic
blood pressure levels, lasts for ≤5 minutes, and is responsive to a 1 g intravenous bolus of calcium chloride and or intravenous boluses of epinephrine (≤100ug) without requiring continuous infusion of vasopressor agents [102]. Severe PRS is defined as the presence of persistent hypotension >30% of the anhepatic level, asystole, significant arrhythmias, and requirement of intraoperative or postoperative vasopressor support [102]. Severe PRS is additionally defined as prolonged (>30 minutes) or recurrent fibrinolysis requiring treatment with antifibrinolytics. The three main categories that contribute to the development of PRS are donor/organ related, recipient related, and procedure related [103]. Prolonged warm ischemia time typically >90 minutes is a procedure related factor that can contribute to the increased risk of developing PRS.

The reality is that there is an interplay between many risk factors that contribute to PRS. In the setting of a technically straightforward transplant with an optimal donor, the new liver begins to produce coagulation factors immediately, and is able to metabolize systemic toxins, thus avoiding PRS and potential primary graft non-function. In the setting of a technically challenging transplant and higher donor risk index organ, the newly implanted liver may have difficulty in initially metabolizing the pooled systemic blood while simultaneously synthesizing necessary coagulation and antithrombotic factors.

During the neohepatic phase, the donor and recipient hepatic arteries are reconstructed with 6-0 to 7-0 Prolene sutures depending on size of the hepatic artery. Various arterial reconstruction techniques can be employed along with different recipient and donor arteries depending on donor and recipient anatomy [104]. Commonly, an end-to-end parachute technique (between donor and recipient common hepatic arteries) is performed. Alternatively, a Carrel patch of donor celiac artery can be anastomosed to the recipient common hepatic artery. A cholecystectomy and bile duct reconstruction are performed, and the abdomen is closed. If technically feasible, an end-to-end bile duct anastomosis is preferred. Alternatively, a Roux-en-Y hepaticojejunostomy can be performed. Intra-abdominal drains are placed at the surgeon’s discretion.

Point-of-care coagulation monitoring with thromboelastography (TEG) or rotational thromboelastometry (ROTEM) has become commonly utilized within LT. Both TEG and ROTEM measure the viscoelastic properties of clot formation via whole blood assay tests that analyze the phases of clot formation [105] and fibrinolysis [106]. Both technologies can measure coagulopathy more accurately than standardized laboratory tests. Additionally, TEG and ROTEM have fast turnaround times. Standard laboratory tests measure coagulation in plasma, are associated with a 40–60-minute delay, and platelet function is not concurrently assessed [107].

Preoperative TEG has been shown prospectively to help predict which patients will require massive transfusion within 24 hours of surgery [108]. Preoperative ROTEM has also shown promise in predicting bleeding risk during LT [109]. A prospectively randomized trial of 28 patients undergoing orthotopic LT was performed utilizing intraoperative TEG compared to standard laboratory measures. Intraoperative TEG monitoring was shown to significantly reduce transfusion rates of plasma (12.8 U vs 12.5 U); however, 3-year survival was not affected [110].
Furthermore, intraoperative use of prothrombin complex and cryoprecipitate guided by ROTEM has shown to result in significantly less RBCs and FFP being transfused [111]. Overall, ROTEM and TEG have shown to help reduce perioperative blood loss and blood transfusions and are rapidly becoming indispensable adjuncts during LT [106].

With the wide adoption of tranexamic acid (TXA) to help reduce bleeding in trauma, there has been interest of adopting the use of TXA during LT. A large systematic review and meta-analysis of liver transplant recipient outcomes comparing the use of antifibrinolytics to placebo found that there was no increased risk for hepatic artery thrombosis, venous thromboembolic events, or perioperative mortality [112]. However, international recommendations advise against the prophylactic use of tranexamic acid [113], unless fibrinolysis is detected clinically or with point-of-care testing. ROTEM has also demonstrated to be helpful in guiding resuscitation in response to hyperfibrinolysis [114].

Postoperative Management After Liver Transplantation

Systemic and renal vascular changes associated with cirrhosis-induced hyperdynamic circulation have been demonstrated to return to normal after LT. However, several authors have also demonstrated that cirrhosis-induced hyperdynamic circulation persists for a long period of time post-LT despite normalization of liver function and portal pressure [115, 116]. Living donor liver transplant (LDLT) recipients with a good postoperative course have been found to have significantly higher portal venous velocity and volume compared to LDLT recipients with graft failure, while no significant differences were observed in absolute cardiac output, cardiac index, blood volume, mean arterial pressure, and hepatic arterial flow [116].

Postoperative LT complications can be divided into acute and chronic and further divided into vascular and nonvascular complications. The rates of the complications include hepatic artery stenosis (2–13%), portal vein stenosis (2–3%), arterial dissection, pseudoaneurysm (most commonly at the hepatic arterial anastomosis), or hepatic artery rupture (0.64%) [117]. Hepatic pseudoaneurysms typically appear in the second to third weeks post-LT with an incidence of 2.5% [118].

Hepatic artery thrombosis (HAT) is the most common acute vascular complication and is considered to be the most devastating as it contributes to bile duct necrosis, graft loss requiring re-transplantation, and overall mortality rates between 27 and 58% [119]. Early HAT is defined as occurring within 1 month of LT and has a higher reported mortality rate compared to late HAT (defined as >1 month post-LT [120]. Early HAT incidence can range from 0 to 12% [117]. A systematic review of 21,822 liver transplants identified 843 cases of early HAT with a mean incidence of 3.9% and without any significant difference between transplant centers worldwide [121]. Of note, this large review defined early HAT as occurring within 2 months of LT. The authors identified that low-volume centers (<30 transplants per year) had a higher incidence of early HAT compared to high-volume centers (5.8% vs 3.2%)
Furthermore, it was demonstrated that pediatric HATs occurred with significantly higher incidence compared to adults (8.3% vs 2.9%) [121]. There is also no significant difference in the incidence of HAT between deceased donor LT (4.6%) compared to LDLT (3.1%) [121]. The median time to diagnosis of early HAT is 6.9 days and of late HAT is 6 months [121, 122].

Risk factors for HAT include increased graft ischemia time, ABO incompatibility, CMV infection, acute rejection, and use of aortohepatic conduit anastomosis, although this can be overcome with experience [123]. Surgical causes for early HAT include retrieval injuries, technical failure, hepatic artery kinking, and small or multiple arteries requiring arterial reconstruction [118]. The type of arterial reconstruction impacts graft function likely secondary to kinking. Long-artery grafts are an independent risk factor for early HAT, and short-graft artery reconstruction is recommended [124].

A systematic review of 19 studies identified that when standard revascularization techniques were not feasible and arterial conduits were utilized, there was an independent increased risk for the development of HAT and increased risk of ischemic cholangiopathy and lower graft survival compared to LT without arterial conduits [125]. Schroering et al. performed a 10-year retrospective analysis of 1145 transplants and identified that nontraditional donor arterial anatomy did not result in any significant difference in HAT or 1-year graft survival [126]. Sixty-eight percent of livers had standard anatomy, 222 donor livers required back table reconstruction, and the most common reconstruction (161 cases) was of the accessory/replaced right hepatic artery to the gastroduodenal artery [126].

Routine early postoperative doppler ultrasonography (US) for the evaluation of HAT has been previously proposed [127], and it is routinely used in many centers for screening for postoperative vascular complications. It is common to obtain postoperative day 1 doppler US to rule out an obvious HAT as one can develop within a few hours of LT. Doppler US is also useful to establish baseline hepatic flows for future comparison. Protocols for postoperative US are variable from center to center. Typical symptoms of early HAT include fever, elevated WBC, elevated transaminases, and possible septic shock; however, patients are often asymptomatic [119]. Doppler US remains as the first-line imaging modality to detect vascular complications as it is relatively quick, inexpensive, and noninvasive [128].

A noncomplicated hepatic arterial anastomosis on US should demonstrate arterial waveforms with swift upstrokes lasting <0.08 seconds, continuous anterograde diastolic flow, and a normal resistive index (RI) of 0.5–0.8 [123]. Transient increases of RI > 0.8 are common within 48–72 hours posttransplantation and are typically due to edema, vasospasm, or the new graft’s initial response to portal hyperperfusion [123]. Increased peak systolic velocities or absent diastolic flow can be seen on US within the first 72 hours and eventually return to normal; however, one must be suspicious of HAT when absent or reversed diastolic flow in combination with low or decreasing peak systolic velocity are present [123, 129]. Additionally, presence of low RI in the initial postoperative period is 100% sensitive for a vascular (arterial, portal, or hepatic) complication [119]. Marin-Gomez et al. identified that low
intraoperative hepatic artery blood flow of 93.3 ml/min was an independent risk factor for early HAT compared to an intraoperative blood flow of 187.7 ml/min without HAT [130].

The development of early HAT will require re-transplantation in approximately 50% of patients [131]. Re-transplantation has traditionally been the primary approach; however, surgical and endovascular revascularization are alternative options. Surgical revascularization has the benefit that the patient does not necessarily need to be re-listed if revascularization is successful. Especially in the current climate of limited organs, surgical revascularization with donor and recipient hepatic artery reconstruction is optimal. Scarinci et al. have reported that when revascularization is performed within the first week of LT, graft salvage approaches 81% [132]. However, successful surgical revascularization rates are variable across the literature [119]. In such situations where surgical revascularization fails, immediate re-listing and re-transplantation are required. In overtly symptomatic patients, with significant hepatic infarction or biliary necrosis, re-transplantation becomes the default primary option. Patients are eligible for immediate re-listing if they are diagnosed with HAT within 7 days of LT, along with an AST ≥ 3000 and/or an INR ≥ 2.5 or arterial pH of ≤ 7.30, venous pH of 7.25, and/or lactate ≥ 4 mmol/L [133].

Endovascular treatments for HAT include intra-arterial thrombolysis and percutaneous transluminal angioplasty with or without stent placement and have been used with increased frequency with some authors reporting high success rates [119, 133]. Endovascular approaches remain somewhat controversial, lack high-quality evidence, and require ongoing further study [117]. Late HAT has a reported incidence rate of 1.7%, and patients typically present with fever, jaundice, and hepatic abscesses [122]. Late HAT with evidence of arterial collateralization should be managed conservatively [118]. However, many patients with late HAT develop ischemic cholangiopathy which requires subsequent re-transplantation [120].

Hepatic artery stenosis (HAS) is defined as narrowing of the hepatic artery by >50% on angiogram with an RI of <0.5 and peak systolic velocity > 400 cm/s [117]. There has been an increasing trend for the management of HAS via interventional procedures. A meta-analysis of case series for HAS was performed by Rostambeigi et al., which identified that percutaneous balloon angioplasty and stent placement have similar success rates (89% and 98%), complications (16% and 19%), arterial patency (76% versus 68%), re-intervention (22% versus 25%), and re-transplantation (20% versus 24%) [134].

Vascular outflow complications include hepatic and IVC thrombosis. Patient symptoms/signs include the need for ongoing diuretic therapy, persistent ascites, or abnormal liver function tests. Persistent ascites has been found to be the most common symptom resulting in investigation for hepatic venous outflow obstruction [135]. Untreated hepatic venous outflow obstruction (HVOO) can lead to graft congestion, portal hypertension, and cirrhosis, which may ultimately compromise graft function and patient survival [136]. HVOO has been reported in about 1–3.5% of patients receiving full-sized grafts and found with increased frequency in re-transplanted patients [135, 137]. Doppler US is the initial radiographic investigation
of choice. Incidence of HVOO for orthotopic LT with partial grafts range from 5 to 13% and 12.5% in LDLT [136]. Early (within 1 month) HVOO is thought to occur secondary to kinking at the donor hepatic vein and recipient suprahepatic IVC anastomosis, technical factors resulting in a narrow anastomosis, or large graft compression of the IVC [137]. Delayed HVOO is related to fibrosis and intimal hyperplasia.

In order to evaluate the incidence of HVOO, a retrospective review of 777 consecutive liver transplants including 695 cadaveric transplants with a mean MELD score of 14, of which 88% underwent piggy back technique was performed [138]. Early hepatic vein outflow obstruction occurred in 1% (7/695) of cases with all occurrences in the piggyback technique with 2 hepatic veins [138]. Two of seven cases were successfully managed medically with diuretics, while five of seven cases required operative cavoplasty [138]. In patients with high-pressure gradients or hepatic vein stenosis at the anastomotic site, hepatic venoplasty alone has been used as the initial management strategy followed by hepatic vein stenting if symptoms or elevated pressure gradient persist [135]. Other centers have successfully performed venoplasty with stenting as a primary option rather than venoplasty alone [137, 139, 140]. Endovascular management for HVOO is preferred over surgical repair because of the increased morbidity and mortality associated with surgical repair [139]. In LDLT recipients diagnosed with early and late HVOO managed with stent placement, patency in the early HVOO group was 76, 46, and 46%, while late HVOO patency rates were 40, 20, and 20% at 1, 3, and 5 years, respectively [141].

Nonvascular complications are further subdivided into biliary complications, graft dysfunction/rejection, infectious, drug toxicity, and increased future risk of malignancy. Biliary complications are the most common complications post-LT, and duct ischemia is closely related to hepatic arterial complications. The biliary system is supplied only by the hepatic arterial system, and arterial anastomotic complications may lead to secondary biliary complications. Common biliary complications include strictures, leaks, stones, bile debris, and ischemia. Bile leaks and strictures occur in 2–25% of cases and comprise the majority of postoperative complications [142]. In a large American data set of 12,803 liver transplants, the incidence of bile duct complications was significantly higher in donation after cardiac death (DCD) recipients (23%) compared to neurologic death donor (NDD) recipients (19%) [143]. Within the same database, DCD recipients required more frequent diagnostic/therapeutic procedures (18.8% vs 14.4%), surgical revision of biliary anastomosis (4.1% vs 2.8%), and re-transplantation (9.1% vs 3.8%) when compared to NDD recipients [143]. A large meta-analysis also identified that biliary complications were significantly increased in DCD recipients compared to NDD recipients (26% versus 16%) [144]. Overall incidence of ischemic cholangiopathy was 16% in DCD recipients compared to 3% in NDD recipients [144].

Early bile leaks are defined as those occurring within 4 weeks of LT and usually occur at the site of the anastomosis. Patients may be asymptomatic or present with nonspecific symptoms such as fever and abdominal/shoulder pain and may develop peritonitis with or without superimposed infection. Elevated bilirubin is usually present along with elevations in lab values (GGT/ALP). Diagnosis can be made
with ultrasound, with CT scan, or with magnetic resonance cholangiopancreatography (MRCP) [144]. Several management options are available. Endoscopic retrograde cholangiopancreatography (ERCP) with or without stent placement is typically utilized. Radiographically guided percutaneous drainage can be effectively used in addition to ERCP to drain a biloma. ERCP has the advantage that it is simultaneously both diagnostic and therapeutic. If the bile duct is reconstructed in an end-to-end fashion, ERCP is technically feasible. When a hepaticojejunostomy has been performed; ERCP is more challenging, requiring a skilled endoscopist, and not always technically possible. If ERCP cannot be performed, or is unable to reach the area of concern, percutaneous transhepatic cholangiography (PTC) and drainage are required. If ERCP is unable to adequately stent or reach a leak at the biliary anastomosis, PTC can be additionally performed to control the leak. If a large biliary anastomotic defect or biliary necrosis is present early in the postoperative period, surgical revision with a redo end-to-end anastomosis, choledochojejunostomy, or hepaticojejunostomy is required. The biliary defect may be too large or degree of the biliary necrosis too significant to preserve enough bile duct length for a redo end-to-end anastomosis.

Bile duct strictures mostly develop at the anastomotic site; however, non-anastomotic strictures may develop and are alternatively known as ischemic type strictures [145]. Non-anastomotic strictures can be caused by microangiopathic factors (prolonged cold/warm ischemia, hemodynamic instability) or secondary to HAT [145]. Extraction of the native recipient liver results in loss of arterial collateral circulation, and the newly implanted donor liver will not have arterial collateral circulation to supply the biliary system. Therefore, its blood supply is dependent on the hepatic arterial anastomosis. It takes approximately 2 weeks for collaterals to start to form. When blood flow is reduced to the biliary system, ischemic strictures may develop anywhere along the bile duct. Ischemic bile duct strictures are typically longer than anastomotic biliary strictures, are present in multiple locations, and are usually found at the hepatic hilum; however, they may be present throughout the intrahepatic biliary system [142].

Periportal edema, residual ascites, or fluid around the peri-hepatic space is expected and usually resolves within a few weeks. Normal postoperative US findings consist of periportal edema, reperfusion edema, and fluid stasis [129]. Periportal edema seen on ultrasound can be mistaken for biliary dilatation and was initially thought to correlate with rejection; this has since been disproved [129]. The incidence of acute graft rejection increases with time. Eighteen percent will experience acute rejection within the first 6 months, and this will increase to 33% by 24 months posttransplant [146].

**Scoring Systems for Transplantation**

The survival outcomes following liver transplant (SOFT) score is based on 4 donors, 13 recipient, and 1 operative factor [34]. It was designed in an attempt to improve organ allocation by avoiding transplantation of organs into patients when predicted
survival is below accepted levels. The SOFT score is composed of two components. There is the pre-allocation score to predict survival outcomes following LT (P-SOFT) and a SOFT score that is used to predict survival posttransplantation [34]. The SOFT score has additional variables with allotted points that can be added or subtracted from the P-SOFT score. The SOFT score can be used by the physician as an adjunct in deciding whether to accept a liver organ by estimating the 3-month postoperative mortality rate compared to a MELD estimated 3-month wait-list mortality rate. The SOFT score is the most accurate predictor of 3-month recipient survival and is also accurate for predicting 1-, 3-, and 5-year post-LT survival [34]. It was determined by the authors that the SOFT score was most accurate based on area under the curve analysis. Furthermore, the SOFT score can be used to improve donor-recipient matching [34].

The balance of risk (BAR) score was developed with a similar goal as the other prognostic scoring systems, and that was to assess post-LT recipient survival. Dutkowski et al. [147] wanted to develop a score based on donor, graft, and recipient factors that were readily available pretransplant and that would have a good correlation to 3-month posttransplant survivorship. Dutkowski et al. used the UNOS database and showed that receiver operator characteristic curves were 0.5, 0.6, 0.6, 0.7, and 0.7 for DRI, MELD, D-MELD, SOFT, and BAR for predicting 3-month patient survival [147]. The BAR score discriminated between survival and mortality with a score of 18. A cited advantage of the BAR score is that its included variables are collected in a standard method internationally and that with less variables compared to other scoring systems, it lends itself to quick and readily accessible calculations [147].

The UCLA group wanted to identify predictors of futility and long-term survival in adult recipients undergoing primary cadaveric orthotopic LT for patients with ESLD and MELD scores >40 [20]. They created a posttransplant futility risk score based entirely on independently verified recipient factors that predicted futility. The variables were MELD score, pretransplant septic shock, cardiac risk, and age-adjusted Charlson comorbidity index [20]. Various calibrated coefficients were added to the included recipient variables. A review of currently available scoring systems with associated variables and pertinent points regarding each scoring system is listed in Table 14.3.

### Donor-Recipient Matching for a Sick Patient

Briceno et al. [151] summarize the historical and current realities of donor-recipient matching based on different organ allocation systems, from patient-based, donor-based, or combined donor-recipient-based policies. The higher the MELD score, the lower the mortality risk for deceased donor transplant recipients compared to wait-list candidates, as mortality was more likely to occur while awaiting LT, rather than from risk of mortality at 1 year posttransplantation [11]. Alternatively, deceased donor transplant recipients with MELD scores <15 had a higher risk of
Table 14.3 Posttransplant morbidity scoring systems

| Scoring system                                      | Incorporated variables                        | Pertinent points                                                                 |
|-----------------------------------------------------|-----------------------------------------------|----------------------------------------------------------------------------------|
| **Donor Risk Index (DRI) [148]**                    | Donor characteristics                         | Donor age > 60 strongest risk factor for graft failure                           |
|                                                     | Age                                           | Split/partial thickness associated with >50% risk of graft failure               |
|                                                     | Height                                        | compared to neurologic death donors                                            |
|                                                     | Race                                          | Recipient factors are not included                                             |
|                                                     | Cause of death (CVA)                          | Poor predictive value for patient survival posttransplantation                 |
|                                                     | Cardiac death                                 | ECD are compared to an optimal reference donor with a DRI of 1.                 |
|                                                     | Partial and split grafts                      |                                                                                  |
|                                                     | Location                                      |                                                                                  |
|                                                     | Cold ischemia time                            |                                                                                  |
| **Survival outcomes following liver transplant (SOFT) score [34]** | Age > 60                                      | Warm ischemia excluded                                                          |
|                                                     | BMI > 35                                      | Overlapping variables                                                           |
|                                                     | One previous transplant                       | Provides a relative risk for 3-month survival                                   |
|                                                     | Two previous transplants                      | <5 points, low risk                                                             |
|                                                     | Previous abdominal surgery                    | 6–15 points, low-moderate risk                                                  |
|                                                     | Albumin <2.0 g/dL                             | 16–35 points, high-moderate risk                                                |
|                                                     | Dialysis prior to transplantation            | 36–40 points, high risk                                                        |
|                                                     | ICU pretransplant                             |                                                                                  |
|                                                     | Admitted to hospital pretransplant            |                                                                                  |
|                                                     | MELD score > 30                               |                                                                                  |
|                                                     | Life support pretransplant                    |                                                                                  |
|                                                     | Encephalopathy                               |                                                                                  |
|                                                     | Portal vein thrombosis                        |                                                                                  |
|                                                     | Ascites pretransplant                         |                                                                                  |
|                                                     | P-SOFT (based on above variables)            |                                                                                  |
|                                                     | Portal bleed 48 h pretransplant               |                                                                                  |
|                                                     | Donor age 10–20 years                         |                                                                                  |
|                                                     | Donor age > 60                                |                                                                                  |
|                                                     | Donor cause of death from CVA                |                                                                                  |
|                                                     | Donor creatinine >1.5 mg/dL                   |                                                                                  |
|                                                     | National allocation                           |                                                                                  |
|                                                     | Cold ischemia time 0–6 hours                  |                                                                                  |
| **BAR score [147]**                                 | Recipient age                                 | Total score out of 27                                                            |
|                                                     | Recipient MELD score                          | Score > 18 considered futile, although this represents only 3% of liver transplants |
|                                                     | Re-transplantation                            | Recipient MELD score strongest predictor of 3-month mortality                   |
|                                                     | Recipient pretransplantation life support     | Less variables than SOFT                                                        |
|                                                     | Cold ischemia time                            | Pretransplant variables removed from score:                                   |
|                                                     | Donor age                                     | Dialysis                                                                       |
|                                                     |                                               | Encephalopathy                                                                  |
|                                                     |                                               | Ascites                                                                         |
|                                                     |                                               | Portal bleeding                                                                 |

(continued)
post-LT mortality at 1 year compared to candidates (with MELD <15) awaiting transplantation. In this analysis, the quality of donor organ was not accounted for. When high MELD score patients receive high-risk or optimal organ donors, there is a survival benefit regardless of the DRI [10]; however, in patients with low MELD scores that received high DRI organs, there is an overall decrease in post-transplant survival [26].

As transplant wait-lists continue to increase along with patients accumulating higher MELD scores and limited organ supply, the use of extended donor criteria has increased. The importance of optimal donor-recipient matching has heightened. Recent data has revealed that 20-year survival for post-LT recipients is significantly influenced by the DRI (≤1.4 and >1.4) and donor age independently (<30 vs ≥30) [152]. It has been suggested that the ideal liver transplant recipient is a young woman with acute liver failure or cholestatic liver disease/autoimmune hepatitis, who has as BMI < 25, normal kidney function, and no dyslipidemia, while the optimal donor organ is <30 years old with an ET-DRI of <1.2 [152]. This optimal match is a rarity in today’s clinical practice, and identifying donors that provide the best match for the sickest first or high MELD priority allocation system is paramount. An ideal match between donor and recipient would ensure that recipient survival and graft survival were optimized, where the probability of death on wait-lists, posttransplant survival, overall cost-effectiveness, and global survival benefit are all accounted for [151]. The question arises, should or shouldn’t a liver be accepted for a particular patient while being cognizant of not just the immediate survival benefit of the particular recipient but also of the factors previously mentioned?

| Table 14.3 (continued) |
|------------------------|
| **D-MELD [149]**       | Recipient MELD score multiplied by donor age | Easily calculated Quick reference for high risk donor/recipient matches Score ranges between 40 and 3400 Score > 1600 found to have worse survival compared to <1600 Donor must be <54 years old for every MELD >30 recipient Patient and graft survival at 4 years |
| **Delta-MELD [150]**   | Total change in MELD points from time of placement on waiting list to transplantation | Does not independently predict mortality after transplantation |
| **UCLA-FRS [20]**      | Recipient MELD score Pre-OLT septic shock Cardiac risk Age-adjusted Charlson comorbidity index ≥6 (CCI) | Entirely based on recipient risk factors Recipient factors predicted futility rather than demographic, donor, or operative factors Cardiac and age-adjusted comorbidities associated with highest risk for futile outcome |

| **Table 14.3 (continued)** |
|-----------------------------|
| **Delta-MELD [150]**       | Total change in MELD points from time of placement on waiting list to transplantation | Does not independently predict mortality after transplantation |
| **UCLA-FRS [20]**          | Recipient MELD score Pre-OLT septic shock Cardiac risk Age-adjusted Charlson comorbidity index ≥6 (CCI) | Entirely based on recipient risk factors Recipient factors predicted futility rather than demographic, donor, or operative factors Cardiac and age-adjusted comorbidities associated with highest risk for futile outcome |
Previously, it was believed that high DRI organs should not be transplanted into patients with high MELD scores [27]. Further study revealed that in patients with high MELD scores, the donor organ quality measured by the DRI did not affect graft or patient outcomes, while in low to intermediate MELD score patients, the DRI did impacts graft/recipient survival [28]. Rana et al. [34] provide recommendations for donor-recipient matching according to recipient MELD score and donor quality as per SOFT score, displayed in Table 14.4.

Rauchfuss et al. [153] reviewed 45 patients who underwent LT with a MELD score of ≥36; their goal was to assess if DRI was associated with 1-year recipient survival post-LT. It was identified that the median duration of waiting time (2 days versus 4 days) was the only significant factor on univariate analysis that differentiated survivors from non-survivors. The overall survival in the group’s study was 69.8% at 1 year. The DRI (median survivors 1.72 vs median non-survivors 1.89), mechanical ventilation status, use of vasopressors, renal replacement therapy prior to LT, or presence of the lethal triad (coagulopathy, hypothermia, acidosis) did not significantly differentiate between survivors and non-survivors [153]. The overall DRI was quite high; however, there was no significant difference between survivors and non-survivors for extended donor criteria. The definition of extended donor criteria included donor age > 65, donor BMI >30, ICU stay >7 days, histologic proven graft steatosis >40%, donor sodium >165 mmol/l, or more than three times increased AST, ALT or bilirubin, donor malignancy history, positive hepatitis serology, drug abuse, sepsis, or meningitis [153].

**Liver Transplantation in Patients with Portal Vein Thrombosis**

Portal vein thrombosis (PVT) is usually diagnosed incidentally in patients with underlying cirrhosis and may affect those with compensated or decompensated cirrhosis. PVT most commonly occurs in patients with cirrhosis with a prevalence of 1–16% [154]. PVT can also occur in patients with hepatocellular carcinoma (HCC) and other hepatobiliary malignancies. Different series report a 2.1–26% incidence of PVT in ESLD patients awaiting LT [155]. More recent data has reported that HCC and cirrhosis carry a 23–28% risk of PVT [156].

Cirrhosis is the clinical manifestation of derangements in the hepatic architecture secondary to fibrosis leading to an increased portal resistance, decreased velocity of blood flow, and subsequent development of collateral venous circulation. Reduced
flow and increased pressure within vessels create stasis and potential for clot formation. PVT in an underlying cirrhotic patient may contribute to further increase in venous pressures, leading to worsening portal hypertension and decreased synthetic liver function [157]. Cirrhotic patients with PVT have an increased association with factor 5 Leiden and prothrombin gene mutations. Mutation in the 20,210 gene has been shown to be an independent risk factor for the development of PVT [158].

Regardless of the underlying etiology of PVT, patients may present with an acute, subacute, or chronic PVT which may result in a partial or complete portal vein occlusion. PVT is further subdivided into benign versus malignant and intrahepatic versus extrahepatic thrombosis [159]. Extrahepatic PVT is exceedingly more common than intrahepatic PVT. For brevity, when discussing PVT, it will be inferred that it is an extrahepatic PVT unless stated otherwise. It is important to distinguish between acute versus chronic PVT and partial versus occlusive thrombus as management strategies, morbidity, and mortality vary accordingly.

Chronic PVT usually presents in an asymptomatic fashion and is incidentally found on imaging performed for other indications or during screening of cirrhotic patients awaiting LT. Chronic PVT in the setting of cirrhosis may eventually lead to accelerated sequela of portal hypertension manifested by ascites, variceal bleeding, ectopic varices, anemia, thrombocytopenia, or splenomegaly [160]. In the setting of a symptomatic PVT, gastrointestinal hemorrhage may be the first sign of underlying portal hypertension. Historically, there was an increased risk of death related to bleeding complications secondary to portal hypertension; however, improvements in prophylactic management of esophageal varices have reduced patient morbidity and mortality [161]. Malignant venous thrombus is diagnosed by an enhancement of the thrombus with direct contiguous extension of the tumor into the portal vein with disruption of the vessel continuity on CT, arterial pulsatile flow on doppler US, or by an increased uptake on PET scan [159]. Patients with malignant PVT are not candidates for LT, and therefore malignant PVT must be distinguished from nonmalignant PVT during the transplant evaluation.

Cirrhosis associated PVT treated with therapeutic LMWH has been shown to be safe and successful with complete or partial recanalization in 60% of patients [162]. Patients need to be continued on LMWH despite image documented recanalization. Patients who demonstrate complete recanalization and stop anticoagulation early have up to 38% re-thrombosis risk [163]. In cirrhotic patients, lifelong anticoagulation may be required to maintain a patent portal vein post recanalization. A small randomized control trial of 70 outpatients with advanced cirrhosis randomized patients to receive 12 months of enoxaparin (dosed at 4000 IU/day) versus no treatment. Patients who received enoxaparin had a significantly lower rate of PVT development (8.8% versus 27.7%) at a 12-month follow-up [164]. An interesting finding of the study was that patients who received enoxaparin had a delayed occurrence of decompensated cirrhosis and improved survival compared to controls.

If a patient with PVT has a contraindication for systemic anticoagulation, a TIPS procedure, if technically feasible, should be considered. The advantages of a TIPS procedure over anticoagulation are a decreased risk of bleeding, possibility of utilizing catheter-based interventions, and risk reduction of recurrent PVT. TIPS has
shown to have a 98% technical success rate for PVT treatment pretransplant, with 92% patency rates until transplant or follow-up, without requiring post TIPS anticoagulation [165]. TIPS can also reduce complications of portal hypertension via portal bypass resulting in improved flow. However, there is high risk of hepatic encephalopathy (27% and 32% at 1 and 3 years) [165]. In a large series of nonmalignant PVT, TIPS resulted in complete recanalization in 57% of patients, 30% reduction in thrombus load, and 13% showed no improvement with an overall technical success rate of 100% [166]. It is important to note that the type of stent (bare versus covered) can impact TIPS dysfunction and the surgeon should be aware of which type of stents are available at their institution. Bare stents have been associated with increased TIPS dysfunction at 1 and 2 years (38% and 85%) compared to covered stents (21% and 29%) [166]. Thornberg et al. have also reported that portal venous recanalization TIPS is technically simpler and easier to perform via transsplenic access compared to transhepatic access [165]. Ultimately, treating PVT is important in reducing the risk of developing an occlusive PVT and also reduces the associated technical risks of PVT and LT.

The development of PVT in patients with ESLD awaiting LT was historically considered an absolute contraindication for proceeding with LT [167]. In the past, LT with surgical management of PVT was associated with increased blood loss, coagulopathy, and mortality [168]. It has been reported that cirrhotic patients undergoing transplant evaluation with occlusive PVT have a significantly lower survival compared to those without occlusive PVT (p = 0.007); and occlusive PVT is in itself an independent risk factor for perioperative death [154]. Identifying which patients with PVT benefit most from LT is paramount in order to minimize postoperative complications and optimize postoperative recovery.

Previously, the identification of PVT in patients awaiting LT was considered a relative indication for adding points to the MELD score in order to transplant patients with PVT earlier. A review of 46,530 patients from the Scientific Registry of Transplant Recipients (SRTR) database showed that the presence or absence of PVT in transplant candidates has no difference on survival while awaiting LT [169]. In the presence of PVT, transplant recipients with MELD scores <12 had significantly inferior postoperative survival with a more than fourfold increase in mortality compared to wait-list mortality. The benefit of LT was seen in MELD scores >13 regardless of presence or absence of PVT [169]. Doenecke et al. [170] retrospectively reviewed 170 liver transplant patients and identified that a MELD score < 15, and presence of PVT was associated with significantly higher perioperative mortality (33%) compared to patients with a patent portal vein (5%). Furthermore, 1-year survival was significantly lower in patients with MELD <15 and PVT compared to patients with a patent portal vein (57% versus 89%). In patients with MELD scores >15, there was no statistically significant difference in mortality between a patent portal vein or presence of PVT. An important observation from these studies is that patients with PVT and MELD scores <13 should not be transplanted. A watchful approach is most beneficial along with medical management of the PVT until MELD score increases to at least >13, which would then provide a survival benefit. If PVT patients with MELD scores <13 are diagnosed with porto-pulmonary hypertension
or hepatopulmonary syndrome, they may receive additional MELD score points and be considered for earlier LT.

Improvements in patient selection for LT, operative techniques, and perioperative management have resulted in PVT becoming a relative contraindication in proceeding with LT [171]. Molmenti et al. reported on 85 cases of PVT that were managed with thromboendovenectomy at the time of LT in comparison with a control group without PVT, and there were no significant differences in 1-, 3- and 6-year patient and graft survival rates between the groups [171]. Gimeno et al. [172] demonstrated that the anhepatic phase and transplant duration were only slightly longer in patients with PVT compared to patients without PVT ($p = 0.28$, $=0.23$). Llado et al. [173] showed that PVT at time of LT is not associated with an increase in overall morbidity and mortality. However, PVT is associated with longer operative times, hospital length of stay, and increased RBC transfusions [173].

Currently, the preferred grading system for PVT was established by Yerdel et al. [174] and has four grades, which are treated differently at the time of surgery.

1. PV minimal or partially thrombosed <50% of the vessel
2. >50% PV occlusion
3. Complete thrombosis of the PV and proximal SMV
4. Complete thrombosis of the PV as well as proximal and distal SMV

Commonly employed surgical techniques that are used for PVT Grades 1–4 are thrombectomy, thromboendovenectomy with venous reconstitution, and interposition of vein grafts. In rare circumstances, with extensive PVT, portocaval hemitransposition has been described. Thrombectomy and its technical variants, interposition grafts, and mesoportal jump grafts are techniques that restore physiologic portal flow. Nonphysiologic technical options are portocaval hemitransposition, renoportal anastomosis, and portal vein arterialization.

Grade 1 and Grade 2 PVT are more common than Grade 3 and 4 PVTs. When technically feasible, the procedure of choice for the management of PVT is considered to be thromboendovenectomy. Grade 1 and 2 PVT can be managed with end-to-end portal vein anastomosis with or without thrombectomy. Grade 1 and 2 PVT repaired with simple thrombectomy, eversion thrombectomy, or improved eversion thrombectomy have been associated with 0% in hospital mortality rate [175]. Furthermore, endovenetomy has been shown to successfully restore portal venous flow in 90% of cases of PVT at the time of LT [154]. It has also been shown that partial PVT patients have a similar incidence of postoperative complications to patients without postoperative PVT [176].

Grade 2 and 3 cases may be amendable to thrombectomy and end-to-end anastomosis; however, an anastomosis at the SMV confluence may be required instead of the proximal portal vein. When the distal SMV is not available for anastomosis, dilated branches of the recipient portal venous system (coronary vein or large collateral vein) may be utilized. Despite intraoperative technical advances in the management of PVT and reported equal survival between PVT and no PVT transplant patients, increased PVT grade is still reportedly associated with worse in hospital mortality, secondarily to increased technical difficulty of successful thrombectomy
techniques [175]. Specifically, Grade 2 or higher PVT has been associated with increased risk of perioperative complications, mortality, and decreased long-term survival [160].

Grade 4 PVT is the most technically challenging for the transplant surgeon. Grade 4 PVT can be operatively managed with anastomosis to the coronary vein or a dilated collateral vein, and eversion thrombectomy procedures are documented to have good outcomes [173, 175]. If the previously mentioned technical options are not feasible, portocaval hemitransposition is an alternative technique that is generally accepted as a last resort. Some authors state that the portocaval hemitransposition technique should be the standard surgical approach for Grade 4 PVT (ref).

Postoperative PVT rate in preoperative complete and partial PVT have reported to be 22.7% and 3.3% with a de novo postoperative PVT rate of 1.3% in patients with a preoperative patent portal vein [176]. Jia et al. [176] in 22 cases of complete PVT and 33 cases of partial PVT most commonly performed a PV reconstruction was an end-to-end PV anastomosis (47 cases), followed by 3 portocaval hemitranspositions, 1 PV to mesenteric vein anastomosis, and 1 PV to renal vein anastomosis, highlighting the use of portocaval hemitransposition as a salvage option in either complete or partial PVT. Additional options are available to the previously mentioned surgical alternatives when portal vein thrombectomy fails to re-establish adequate portal vein flow for extensive PVT. Quintini et al. [177] describe renoportal bypass using a venous conduit from the recipient renal vein anastomosed to donor portal vein. The left renal vein is dissected with a caudal mobilization of the soft tissue anterior to the inferior vena cava, until the left renal vein is identified at the insertion of the IVC. A caveat of the renoportal bypass procedure is that its success is also dependent on the presence of a patent splenorenal shunt.

Regardless of PVT grade, re-establishing physiological portal venous flow has a significant impact on reducing patient morbidity. Hibi et al. [178] retrospectively reviewed a large cohort examining 174 patients with PVT (48% occlusive, 52% partial thromboses) at the time of transplantation. They identified that 149 PVT patients had physiological portal inflow re-established and there was no significant difference in survival between patients with re-established physiological portal inflow compared to patients without PVT. Thrombectomy was performed in 123 cases, while 16 patients received interpositional vein grafts, and 10 patients underwent mesoportal jump grafts. The subsequent challenge is improving outcomes in PVT patients in whom physiological portal inflow cannot be re-established. The same study identified that when physiological portal inflow was not re-established, there was a significant increase in the incidence of re-thrombosis, gastrointestinal bleeding, and worse 10-year overall survival [178]. In the nonphysiologic PVT group, 18 underwent cavoportal hemitranspositions, 6 renoportal anastomoses, and 1 portal arterialization procedures.

In adults, post-orthotopic LT portal venous complications (stenosis or thrombus) occur at a rate of approximately 3%. In the pediatric population, portal vein complications are higher at approximately 8%. The increased complication rate in this population is secondary to the increased technical challenge of a shorter portal vein, the use of living-related donors, and split LT [179]. Generally, portal vein stenosis
occurs at the anastomotic site secondarily to donor/recipient portal vein diameter mismatch [162, 179]. Portal vein stenosis may occur in the immediate postoperative period or can be detected during long-term follow-up. Patients may be asymptomatic or present with signs of portal hypertension, similar to pretransplant PVT presentation.

Management of posttransplant portal vein stenosis differs from the management of posttransplant PVT. Portal vein stenosis management is dependent on whether the stenosis is deemed to be clinically significant or not. In asymptomatic patients with normal hepatic function, periodic observation with ultrasound has been described. However, in patients where portal venous stenosis is potentially contributing to worsening, portal hypertension intervention is required [179]. Interventional percutaneous portal vein dilatation with or without stent placement can be performed. Funaki et al. [180] and Shibata et al. [181] describe treatment of portal venous stenosis with balloon dilatation and stent insertion if pre- and post-portal vein dilatation pressure gradient is >5 mmHg or >3 mm Hg. Funaki et al. have had high success with interventional venoplasty and have eliminated the need for surgical revision, portacaval shunting or re-transplantation [180]. Management of PVT post-LT may differ between various institutions. Experience with percutaneous vein thrombolysis is limited, and few case reports have been published. For significant postoperative PVT that is not amendable to anticoagulation, portal vein angioplasty (with or without stent placement) remains as a first-line option, followed by TIPS, or re-transplantation, as a last resort.

**Impact of Hepatic Flows in Liver Transplant**

The liver weighs approximately 1.2–1.6 kg and is 2.5% of a human’s total body weight, yet receives 25% of the cardiac output [182, 183]. Total hepatic flow ranges between 800 and 1200 mL/min [184]. The hepatic inflow is supplied both by the hepatic artery and portal vein. Twenty-five percent of the total hepatic flow comes from the hepatic artery, which provides 30–50% of hepatic oxygen requirements [185]. Seventy-five percent of the total hepatic flow is provided by the portal vein, which provides 50–70% of hepatic nutritional requirements [186]. Interplay between the portal and hepatic inflow has significant impact on hepatic regeneration [187]. This is especially true after orthotopic LT and even more so for LDLT. Forty percent of the hepatic blood is within large vessels, while 60% is held within the hepatic sinusoids. Hepatic sinusoids are very compliant and can accommodate a large volume of blood so that portal venous flow can be increased or decreased without disruption to the portal venous pressure in healthy livers [185]. Significant differences can be seen in portal venous blood flow between non-cirrhotic and cirrhotic patients especially if extensive portal hypertension or PVT is present. Cirrhotic-induced vascular changes impact intraoperative decision-making on whether to perform standard donor to recipient vascular anastomosis versus a modified restoration of physiologic or nonphysiologic inflow. The goal is to provide
optimal blood flow and tissue perfusion that are required for the metabolic activity of the liver [185].

The hepatic arterial system is a high-pressure, high-resistance system with an average flow of 400 mL/min that is controlled by an intrinsic autoregulatory system [188]. Norepinephrine and angiotensin can cause hepatic artery vasoconstriction without affecting the portal vein flow. Their effects can be reversed with high doses of intra-arterial adenosine [189], a well-known vasodilator. The main portal vein provides 50–70% of the liver’s oxygen requirements with a flow of 700–850 mL/min and portal pressure ranging between 5 and 10 mm Hg in healthy subjects. The high-resistance arterial system ends at hepatic sinusoids and transitions to the portal venous system via sinusoidal capillaries [190]. Under normal conditions, the portal venous system is a low-pressure, low-resistance system. It is affected by venous drainage from visceral organs, regulated by splanchnic arteriole constriction and intrahepatic vascular resistance. The hepatic artery and portal vein flows in healthy individuals are typically proportional; however, in patients with cirrhosis, it is observed that the portal vein and hepatic artery flows are inversely related to each other [191].

Adenosine is secreted at a constant rate and is equal between the hepatic artery and portal vein [184]. When portal vein inflow increases, it causes the adenosine to be washed away with a resultant decrease in hepatic artery flow mediated by hepatic artery vasoconstriction [189]. When portal perfusion decreases via impeded or diverted portal inflow, the liver triggers adenosine to locally accumulate. Adenosine induces hepatic artery vasodilatation and increases hepatic artery blood flow in order to compensate for the reduced portal venous flow. In addition to adenosine, local nitric oxide, carbon monoxide, and a gaseous mediator known as H2S have been found to change in concentration depending on the portal venous pressure [189]. The ability of the hepatic artery to vasodilate in response to changes in portal pressure is an intrinsic autoregulatory mechanism of the liver that is known as the hepatic artery buffer response (HABR) [184]. Initial clamping of the splenic artery has shown to cause an increase in hepatic artery velocity followed by a quick and maintained decreased portal venous velocity [192]. Subsequent clamping of the splenic vein induces a significantly quick and maintained decrease in the portal venous flow, with an eventual increase in hepatic arterial flow [192]. The increase in hepatic arterial blood flow is able to compensate for up to a 25–60% reduction in portal flow [189]. The goal of the HABR is to maintain adequate oxygen supply to tissues and minimize the impact of portal venous flow changes on hepatic clearance [184].

Prior to the HABR theory, it was believed that the splenic artery was diverting blood away from the hepatic artery, and this was termed the splenic steal syndrome [193]. Splenic steal syndrome cannot be diagnosed in the presence of HAT, HAS, arterial kinking, or other hepatic arterial abnormality that may impede its flow [193]. The HABR has been shown to remain intact with human LT [190]. In partial donor liver grafts, there is a blunted HABR response where the hepatic artery inflow remains depressed compared to what would be expected in a whole-sized graft [190].
Doppler ultrasound is a very useful imaging modality to identify physiologic and pathophysiologic hepatic flow. Normal portal vein waveform normally shows gentle undulations with hepatopetal flow [194]. The normal waveform within the hepatic veins is triphasic with two hepatofugal phases related to the atrial and ventricular diastole. Cirrhotic changes in the liver can cause a large reduction in the visualization of hepatic veins that alters the normal waveform. The normal triphasic waveform can be replaced with a monophasic pattern that indicates high portal pressures [194]. Hepatic artery flows can be indirectly measured by the pulsatility index or estimations of the resistive index on doppler ultrasound which approximates the hepatic artery flow [191]. Portal inflow becomes partially reversed, and this is known at hepatofugal flow. Hepatofugal flow of the main portal vein is a known marker of portal hypertension, and it has been identified that a threshold velocity of 11 cm/s in the right portal vein and left portal vein velocity of <8 cm/s are associated with the development of main portal vein hepatofugal flow [195].

In cirrhosis, underlying fibrosis-induced architectural changes result in alterations of hepatic microvasculature, hepatic sinusoids, reduced blood supply, and increased total hepatic vascular resistance. Due to the increased hepatic vascular resistance, the intrahepatic endothelial cells produce less nitric oxide resulting in portal hypertension (mean intraluminal portal pressure > 12 mm Hg). In response, the extrahepatic mesenteric vascular beds cause progressive vasodilation of splanchnic vasculature secondary to increased release of nitric oxide. At baseline, the HABR in cirrhotic patients is continuously active; however, hepatic artery flow changes are blunted in response to sudden changes in portal venous flow [191, 196]. In a cirrhotic patient prior to LT, portal flow is approximately 1 L/min [188, 197]. The porto-splanchnic system attempts to redistribute the increased portal inflow; however, because of cirrhosis-induced fibrosis and increased intrahepatic vascular resistance, the liver is unable to accommodate for the increased incoming portal inflow. Preexisting and/or newly formed venous collaterals receive redistribution of hepatofugal portal flow. This leads to varices and further subsequent vascular remodeling with an overall reduction in portal venous blood flow, increased hepatic venous resistance, systemic hyperdynamic circulation, and increased cardiac output [198]. To compensate for increased intrahepatic resistance, the HABR increases hepatic arterial flow by a reduction in hepatic arterial resistance.

The hepatic artery can induce a compensatory vasoconstriction reducing arterial blood flow in response to portal hyperperfusion and therefore leads to a high resistivity index (RI). Unlike the portal vein patterns, the hepatic artery and the superior mesenteric artery RI do not correlate with the stage of cirrhosis [199]. Hyperdynamic cardiovascular changes can lead to significant obstacles at the time of LT. Sudden reduction in vascular preload and impaired cardiac contractility can impair cardiac output, while in the postoperative period, hypovolemia and hypervolemia can negatively impact cardiac contractility [200]. Hyperdynamic pretransplant cirrhotic pathophysiology persists posttransplantation for months to years, regardless of the underlying etiology of cirrhosis [200]. In patients with underlying viral cirrhosis, there is a rapid improvement with reduced cardiac output and increased systemic vascular resistance that is not present in alcoholic cirrhosis [200].
Obtaining optimal intraoperative hepatic artery and portal vein blood flow is necessary for a successful liver transplant in the short- and long term. However, optimal flows for the hepatic artery and portal vein are still unknown without strong quality evidence. Prior to LT, in the cirrhotic liver, portal flow is approximately 1–2 L/min [188, 197]. Mean hepatic artery flow has a range from 268 to 584 ml/min, with a resultant cardiac output of 10 L/min. Spitzer et al. found that for full donor implanted grafts, a minimum hepatic artery flow of 250 ml/min is required for improved patient survival; however, flows of >400 ml/min are optimal [201].

Intraoperatively, different presentations of altered hepatic flow may present, namely, portal vein enlargement and splenomegaly, without significant collateral formation or reduced portal vein size with massive collateralization [185]. In the setting of a large portal vein without significant collateralization, improving portal venous and hepatic flow can be achieved with either splenectomy or splenic artery ligation [185]. Alternatively, if the portal vein is smaller than expected with large collateralization, spontaneous splenorenal shunting is likely to have occurred. Some authors report that ligation of major collaterals when portal venous flow is <1 L/min may help with preventing portal hypoperfusion [185]. Ligating large collateral coronary veins greater than 1 cm is thought to increase PV flow by 55–140%, depending on the size of the varix [202]. Common veins to ligate are the coronary vein, inferior mesenteric vein, gastroepiploic vein, splenorenal shunt, and retroperitoneal varices. Large splenorenal shunts can be embolized via percutaneous methods or via intraoperative ligation of the left renal vein. The main causes of decreased portal flow are unrecognized portal mesenteric/splenic vein thrombosis, inadequate portal vein thromboendvenectomy, or large portosystemic collaterals [188]. Once a new liver graft has been transplanted, a lower portal resistance within the new graft allows for improved portal flow, which has been measured to increase to 1.8–2.8 L/min after implantation. Minimum portal vein flow should be >1 L as portal vein flow >1 L mL/min is associated with improved graft survival at 30, 60, and 365 days post-LT in the deceased donor transplantation [188, 197, 201].

Decreased intraoperative hepatic artery flows are thought to be primarily due to technical issues with the anastomosis. However, arterial steal syndrome, celiac artery stenosis, or hypoperfusion secondary to under-resuscitation can contribute to decreased hepatic artery flows. Additionally, mechanical ventilation, hypercarbia, positive end expiratory pressure, hypotension, hemorrhage, and hypoxemia are other intraoperative factors that may reduce hepatic artery flow [203]. In patients who develop hepatic artery strictures, there are significantly lower intraoperative arterial and portal vein flows compared to patients who do not develop hepatic arterial strictures [204]. Low hepatic artery RI after deceased donor LT can be attributed to surgical edema, hepatic artery stenosis, severe aorto-cesiac atherosclerotic disease, arteriovenous or arterial biliary fistula formation, hepatic vein, or portal vein thrombosis [199]. The hepatic artery is the sole blood supply to the bile duct. This is supported by evidence that lower measured hepatic flows have been associated with higher rates of biliary complications after LT [205]. Therefore, ensuring proper hepatic artery flow is imperative to obtaining optimal biliary anastomotic outcomes.
Dealing with Portosystemic Shunts to Prevent Portal Vein Steal

TIPS is primarily indicated in patients with refractory ascites and variceal hemorrhage, with less frequent indications being PVT and HVOO and can be utilized as a bridge to LT [206]. In 2015, a Consensus Conference on TIPS was held to provide recommendations for proper evaluation, technical considerations, patient selection, follow-up, contraindications, and management of complications [207]. TIPS placement as a bridge to LT may result in technical difficulties during the transplant with the shunt extending into the portal vein, hepatic vein, or right atrium; however, TIPS has not shown to have any significant negative impact on graft or patient survival [208]. Even in the setting of cirrhosis complicated by nonmalignant PVT, TIPS is technically feasible and is not associated with increased procedure related complications, stent occlusion, or mortality [166]. TIPS can be used to maintain and improve patency of the portal venous system and reduce the re-occurrence of PVT. It has also been used to decrease the effect of mesosystemic collaterals and shunting of blood away from the liver [166, 209].

Patients with portal hypertension and advanced cirrhosis have increased resistance to portal inflow and develop portosystemic shunts; as a result, blood flow is shunted away from the portal vein and liver via the mesosystemic collaterals, otherwise known as hepatofugal flow [210]. Splenorenal shunts form between the splenic and renal veins and are an example of such spontaneous mesosystemic collateral development. Portal steal syndrome develops after LT when the mesosystemic collaterals persist and continually divert flow away from the newly implanted graft [211]. After full-sized cadaveric orthotopic LT, hepatofugal flow usually resolves, portal vein flow becomes hepatopedal and results in a decreased intrahepatic resistance [210]. However, hepatofugal flow may only slightly decrease or persist post-LT, especially in the setting of partial graft transplantation, and contribute to the development of portal steal syndrome [212]. Hyperdynamic spontaneous portosystemic shunts are present in up to 19% of portal hypertensive patients awaiting LT [210]. The higher the flow from the splenic vein into the renal vein the greater likelihood of significant blood flow diverted away from the liver [209]. When a portosystemic shunt persists post-LT, it may reduce portal inflow/portal venous pressure and impact early hepatic regeneration and harm the new graft [212, 213]. This especially applies in small-for-size grafts after LDLT [211]. Risk factors for recipient portal steal phenomenon include portal hypertension with large varices and natural shunts, chronic liver failure, macrosteatosis, low liver donor mass, donation after cardiac death with prolonged warm ischemia time and receiving a LDLT [210].

It is imperative to detect portal flow steal as early as possible and to manage accordingly, to ensure survival of the newly transplanted liver graft. Large spontaneous splenorenal shunts (> 10 mm in diameter) have been shown to occur in 6.6% of adult LDLTs [211]. Splenorenal shunts <10 mm in diameter are thought to not require intervention as portal pressures post-LT normalizes, and the shunt will eventually collapse [209]. Lee et al. describe their technique of left renal vein...
(LRV) ligation in 44 patients with large splenorenal shunt for portal steal syndrome during partial graft LDLT [211]. At the time of LT, intraoperative portal flow assessment of the ligated portal vein was performed when LRV was unclamped and subsequently clamped. If a large difference in portal vein flow was observed during LRV clamping, then ligation of the LRV was performed prior to hepatic arterial construction. The authors report that all 44 patients recovered well without re-transplantation at a median follow-up of 17 months, with 1 patient passing away secondarily to HCC [211].

In the presence of large spontaneous splenorenal shunts, Castillo et al. have used a portal vein flow threshold (after reperfusion) of \( \leq 1200 \text{ ml/minute} \) to perform LRV ligation, which successfully increased portal flow post-ligation without any consequence to renal function [214]. Tang et al. summarize eight case series of LRV ligation with Lee et al. having the largest series of LRV ligation to date [215]. A patent portal vein is required to proceed with LRV ligation which has been demonstrated to improve portal vein blood flow. This should not be performed in unresectable PVT, portal vein stenosis, or with large portal vein mismatch between donor and recipient [215]. Although LRV ligation has been shown to be safe and effective for dealing with portal vein steal syndrome, definitive consensus indications cannot be made based on size of splenorenal shunts or threshold portal vein flows. Larger multicenter prospective studies are required.

In 26 patients with hepatofugal flow detected on preoperative doppler US or weak flow identified at the time of transplant, direct ligation of large splenorenal shunts was performed intraoperatively with a 7.7% major complication rate and 96.2% survival rate [216]. Eleven of the 26 patients with splenorenal shunts had a preexisting PVT and underwent PV thrombectomy. In contrast to LRV ligation, PVT is not a contraindication for ligating the entire shunt. Splenectomy is an alternative option to ligation of the LRV at the time of LT; however, there is an increased risk of PVT, sepsis, and bleeding [215].

**Use of Live Donors in Sick Patients and Impact of Portal Hypertension on Small-for-Size Syndrome**

There is a universal shortage of available organs to meet demand of patients requiring transplantation. Currently, live donor liver transplantation (LDLT) comprises <5% of all liver transplants performed in the United States [217]. In an attempt to reduce LT wait times and increase the organ pool, LDLT was introduced as an alternative to cadaveric transplantation. LDLT was initially performed within the pediatric population; however, currently LDLT has been implemented for adult LT in high-volume centers. The left hepatic lobe has traditionally been used in the pediatric population for an appropriate donor to recipient size match, accounting for the smaller-sized pediatric population. In adults, left lobe implantation was initially utilized; however, initial results were poor due to small-for-size syndrome (SFSS) and early graft dysfunction. In the late 1990s, adult right hepatic lobe LDLT was
Increasingly utilized in order to circumvent SFSS [218]. Recent studies have shown that left hepatic lobe donation is associated with favorable recipient and donor outcomes compared to right hepatic lobe LDLT [217]. Despite this, right LDLT remains the most commonly utilized lobe in adult LDLT due to the ability of the right lobe to provide consistently more reliable hepatic mass [219].

A major limitation for LDLT is the potential for donor death and postoperative donor complications. The risk of donor death from live liver donation (90 days within surgery) is reported to be 1.7 per 1000 donors (0.17%), which is in keeping with living kidney donor rates [220]. Minor donor complications are reported to occur in approximately 27% of donors, with the most common complications being biliary leaks (9%), bacterial infections (12%), and incisional hernias (6%) [221]. Several studies have shown that donor outcomes with left lobe LDLT is associated with lower complication rates, lower rates of serious complications, and identical 1-, 5-, 10-year recipient survival compared to right lobe LDLT [222–224]. Although donors are associated to have increased postoperative morbidity and mortality, in high-volume centers, donors are able to enjoy good postoperative health and return to preoperative baseline without serious complications [225].

Recipient LDLT complications arise from the donor graft having a reduced hepatic reserve and receiving portal flows that are higher than the donor graft would have received in its original state prior to LT; that would normally be reserved for a whole liver. The most pronounced hemodynamic changes are an increase in portal perfusion rate and cardiac output of the recipient secondary to the effects of cirrhosis [226]. Typically, a whole transplanted liver has a large vascular bed of hepatic sinusoids to accommodate for the increased portal flow and cardiac output [218]. The liver compensates for the increased portal vein flow and cardiac output by activating the HABR, which reduces hepatic artery inflow. The live partial donor graft must manage the hyperdynamic portal circulation secondary to high portal flow immediately after LT. With LDLT it is believed that within minutes of reperfusion, portal hyperperfusion can cause shear stress to hepatocytes, sinusoidal congestion, and hemorrhagic necrosis of peri-sinusoidal hepatocytes [227, 228].

In small-for-size syndrome (SFSS), a donor graft is significantly reduced in size and portal hyperperfusion in conjunction with a smaller graft’s high portal resistance can cause further reduction of hepatic artery inflow via the HABR and resultant de-arteriolization [229]. Doppler studies have shown that hepatic artery vasoconstriction in response to portal hyperperfusion and an exaggerated HABR produce a high resistive index with poor arterial perfusion [199]. Additionally, excessive portal flow can lead to oxidative stress thereby activating the inflammatory cascade leading to further hepatocyte damage [230]. The major concern is for graft dysfunction and secondary biliary complications. The symptoms of SFSS manifest as a pattern of liver dysfunction with associated portal hypertension, diminished arterial inflow, delayed synthetic function, and prolonged cholestasis. In advanced cases of SFSS, patients can clinically decompensate with the development of sepsis, encephalopathy, and death [231]. SFSS is typically thought to occur when the donor graft to recipient weight ratio (GRWR) is <0.8% during the first postoperative week after excluding other causes of graft dysfunction [232].
However, studies have identified that GRWR of 0.6 in LDLT is safe [233, 234]. Others have shown that GRWR of 0.6% and 0.85% is safe in Child Pugh Class A recipients, while Child Pugh Class B and C recipients require GRWR >0.85% for appropriate outcomes [235]. The International Liver Transplantation Society Living Donor Liver Transplant Recipient Guidelines state that the safety limit for minimum GRWR can be less than 0.8% in the setting of improved center experience and patient selection; however, most centers consider GRWR of 0.8% as the lower limit [219].

Intraoperative doppler ultrasonography should be used post-hepatic arterial reconstruction to assess hepatic artery flow and portal vein flow [199]. Portal venous pressure has been considered the most important hemodynamic factor influencing the functional status of the liver and graft regeneration post-LT [199]. It has been demonstrated that portal venous pressure < 15 mm Hg results in improved 2-year survival compared to patients with portal venous pressures >15 mmHg [236]. Wu et al. have demonstrated that high portal venous flow was well-tolerated by right LDLT recipients postoperatively if initial portal pressure was <23 mm Hg and the postreperfusion portal venous pressure was <15 mm Hg [213]. Furthermore, when initial portal venous pressure is ≥23 mmHg, and after reperfusion ≥15 mmHg, patients developed significantly more ascites compared to patients with lower portal venous pressures [213]. Optimal portal venous flows and hepatic arterial inflow remain a topic of debate, dependent on right or left LDLT and the true impact of HABR [185, 188]. It has been shown that portal venous flows <180 ml/min/100 g of liver weight (LW) leads to lower survival [237] and experimental models have supported that optimal outcomes occur with portal venous flows <260 ml/min/100 g LW [238]. It is believed that in order to avoid SFSS, portal venous flows of <260 mL min per 100 g LW are recommended [239] and graft inflow modulation techniques should be employed if the portal venous flow is >250 ml/min/100 g LW [240].

Several techniques have been described to decrease or reduce the impact of SFSS via modulation of graft inflow [241] when portal venous pressures exceed 15 mmHg. Splenic artery ligation [242] is usually the first step in portal flow modulation; however, splenectomy [243], portacaval, mesocaval, and splenorenal shunts are alternative options. Splenic artery ligation reduces portal vein flow by 30% [240] by reducing resistance of the distal hepatic artery and subsequently reducing flow in the splenic circulation. The net effect of splenic artery ligation/embolization results in promotion of liver regeneration and overcoming the effects of portal hypertension and portal hyperperfusion [244]. If elevated portal pressures are identified postoperatively, splenic artery embolization can be performed via interventional radiological methods. Splenectomy is potentially life-threatening, and if splenic artery ligation is technically feasible, it should be a primary management option. Portocaval shunts are believed to be beneficial when lower portal venous flows of 190/mL/ min/100 g LW are present compared to higher flows of 401 mL/min/100 g [245].

A group from Taiwan proposed a flowchart for when to perform graft inflow modulation according to the portal venous pressure and portal venous flows which is briefly described here; however, it is yet to be validated [185]. The group
performed splenectomy in the setting of PVF ≥ 250 mL/min/100 g LW, PVP ≥ 20 mmHg, and without outflow obstruction, or if PVF was ≤100 mL/min/100 g LW, PVP was 15–20 mmHg; hepatic arterial inflow (HAF) was <100 mL/min without anastomotic error. No graft inflow modifications were made if the PVF was ≥250 mL/min/100 g LW and the PVP was <15 mmHg or if PVF was ≥250 mL/min/100 g LW, PVP was 15–20 mmHg, and the HAF was >100 mL/min. International recommendations (class 1, level b) for preventing/treating graft injury and SFSS are to monitor the portal vein/hepatic artery hemodynamics and to use portal inflow modulation techniques [219].

In 2002, the New York State Committee on Quality Improvement recommended that patients awaiting LT with MELD scores >25 should not undergo LDLT [246]. However, LDLT has been demonstrated to have similar postoperative complication rates and survival outcomes compared to DDLT [146, 247]. An adult-to-adult LDLT cohort multicenter retrospective study reported a 13.2% graft failure rate in 385 ALDLT recipients in the first 90 days [248]. The group identified that older recipient age and length of cold ischemia were significant predictors of graft failure, while individual center experience greater than 20 ALDLT was associated with lower risk of graft failure. Also, recipient MELD score was not a significant predictor of graft failure, but this sub analysis was limited to a small percentage (4%) of patients with MELD scores >30. [248]. The same group reported a 90-day and 1-year recipient survival of 94% and 89%, respectively. This was a seminal paper, as this was the first multicenter study of donor and recipient LDLT outcomes. A follow-up study identified that adjusted long-term mortality risk between LDLT and DDT was similar (for recipient gender, age, diagnosis, dialysis, MELD, and donor age) [249].

With persistent limited access to organs and growing evidence identifying equivalent outcomes between LDLT and DDLT, focus should be directed to LDLT for patients with high MELD scores and sick patients awaiting LT. High MELD patients awaiting LT have a high wait-list mortality and, as discussed previously, demonstrate significant benefit from transplantation. If deceased donor organ is not available for sick/high MELD patients, consideration should be made to utilize LDLT. However, ethical issues arise regarding the benefit risk ratio for donors undergoing a significant life-transforming event for a potentially futile recipient transplant outcome. In 2006, at the Vancouver Forum on the Care of the Live Organ Donor LDLT was deemed appropriate for acutely ill and sick transplant candidates [250]. However, LDLT in patients with MELD scores >25 remains controversial. It has been shown that in patients with MELD scores >20 undergoing LDLT, preoperative renal dysfunction, severe hypoalbuminemia, and massive intraoperative RBC transfusion are independent risk factors for in-hospital mortality. In recipients with two or more risk factors, 3-month survival was 25% [251].

Recent 5-year recipient LDLT survival has been shown to be similar to DDLT among patients with MELD scores <20, and it has been postulated that LDLT is underutilized in patients with MELD scores above 20 [252]. Feng summarizes the findings of several authors from both Eastern and Western transplant centers that have demonstrated good survival in patients with elevated MELD scores undergoing LDLT [253]. Selzner et al. in a large series compared outcomes in patients with
MELD scores $<25$ and $>25$ in 271 consecutive adult-to-adult right lobe LDLT [246]. They demonstrated that there was no significant difference in the overall complication rate within 3 months of LT between MELD $<25$ and MELD $>25$ recipients ($51\%$ versus $45\%$, $p = 0.28$). Graft survival between MELD $<25$ and MELD $>25$ was not significantly different at 1 year (92\% versus 83\%), 3 years (86\% versus 80\%) and 5 years (78\% versus 80\%), and patient survival was similar between groups at 1 year (92\% versus 83\%), 3 years (86\% versus 83\%), and 5 years (82\% versus 83\%) [246]. Kaudo et al. have also shown that overall recipient patient survival did differ between patients with MELD scores $<25$ and $\geq 25$ who underwent LDLT [254]. Liu et al. found that LDLT for patients with acute on chronic HBV with mean MELD scores of 36 had similar patient survival compared to elective LDLT in patients with mean MELD scores of 17.8 with a median follow-up of 23 months (88\% versus 84\%) [255]. In 2013, Chok et al. displayed similar 1- and 5-year LDLT recipient survival for MELD $\geq 25$ (95.9\% and 93.2\%) compared to MELD $<25$ (96.9\% and 95.3\%) [256].

For many of these studies, right hepatic lobe LDLT was utilized more often than left lobe LDLT, which highlights the general preference for right hepatic lobe donation, especially in the setting of sick and high MELD score patients. In experienced LDLT centers, transplantation of a high MELD recipient is technically feasible and is associated with good outcomes. With continued education, discussion, and supportive data, hopefully LDLT can aid in the challenge of tackling the sickest patients first and can help decrease the shortage of available organs.

References

1. Wiesner R, Edwards E, Freeman R, et al. Model for end-stage liver disease (MELD) and allocation of donor livers. Gastroenterology. 2003;124(1):91–6. PubMed PMID: 12512033. Epub 2003/01/04.
2. Freeman RB Jr, Wiesner RH, Harper A, et al. The new liver allocation system: moving toward evidence-based transplantation policy. Liver Transpl. 2002;8(9):851–8. PubMed PMID: 12200791. Epub 2002/08/30.
3. Malinchoc M, Kamath PS, Gordon FD, et al. A model to predict poor survival in patients undergoing transjugular intrahepatic portosystemic shunts. Hepatology. 2000;31(4):864–71. PubMed PMID: 10733541. Epub 2000/03/25.
4. Merion RM, Wolfe RA, Dykstra DM, et al. Longitudinal assessment of mortality risk among candidates for liver transplantation. Liver Transpl. 2003;9(1):12–8. PubMed PMID: 12514767. Epub 2003/01/07.
5. Biggins SW, Bambha K. MELD-based liver allocation: who is underserved? Semin Liver Dis. 2006;26(3):211–20. PubMed PMID: 16850370. Epub 2006/07/20
6. Bernardi M, Gitto S, Biselli M. The MELD score in patients awaiting liver transplant: strengths and weaknesses. J Hepatol. 2011;54(6):1297–306. PubMed PMID: 21145851. Epub 2010/12/15.
7. Sharma P, Schauble DE, Goodrich NP, Merion RM. Serum sodium and survival benefit of liver transplantation. Liver Transpl. 2015;21(3):308–13. PubMed PMID: 25504743. Pubmed Central PMCID: PMC4354811. Epub 2014/12/17.
8. Merion RM. When is a patient too well and when is a patient too sick for a liver transplant? Liver Transpl. 2004;10(10 Suppl 2):S69–73. PubMed PMID: 15382215. Epub 2004/09/24.
9. Farkas S, Hackl C, Schlitt HJ. Overview of the indications and contraindications for liver transplantation. Cold Spring Harb Perspect Med. 2014;4(5). PubMed PMID: 24789874. Pubmed Central PMCID: PMC3996378. Epub 2014/05/03.

10. Schaubel DE, Guidinger MK, Biggins SW, et al. Survival benefit-based deceased-donor liver allocation. Am J Transplant. 2009;9(4 Pt 2):970–81. PubMed PMID: 19341419. Pubmed Central PMCID: PMC2895923. Epub 2009/04/04.

11. Merion RM, Schaubel DE, Dykstra DM, et al. The survival benefit of liver transplantation. Am J Transplant. 2005;5(2):307–13. PubMed PMID: 15643990. Epub 2005/01/13.

12. Desai NM, Mange KC, Crawford MD, et al. Predicting outcome after liver transplantation: utility of the model for end-stage liver disease and a newly derived discrimination function. Transplantation. 2004;77(1):99–106. PubMed PMID: 14724442. Epub 2004/01/16.

13. Habib S, Berk B, Chang CC, et al. MELD and prediction of post-liver transplantation survival. Liver Transpl. 2006;12(3):440–7. PubMed PMID: 16498643. Epub 2006/02/25.

14. Murken DR, Peng AW, Aufhauser DD Jr, et al. Same policy, different impact: center-level effects of share 35 liver allocation. Liver Transpl. 2017;23(6):741–50. PubMed PMID: 28407441. Epub 2017/04/14.

15. Karvellas CJ, Lescot T, Goldberg P, et al. Liver transplantation in the critically ill: a multicenter Canadian retrospective cohort study. Crit Care. 2013;17(1):R28. PubMed PMID: 23394270. Pubmed Central PMCID: PMC4054984. Epub 2013/02/12.

16. Aloia TA, Knight R, Gaber AO, et al. Analysis of liver transplant outcomes for United Network for Organ Sharing recipients 60 years old or older identifies multiple model for end-stage liver disease-independent prognostic factors. Liver Transpl. 2010;16(8):950–9. PubMed PMID: 20589647. Epub 2010/07/01.

17. Asrani SK, Saracino G, O’Leary JG, et al. Recipient characteristics and morbidity and mortality after liver transplantation. J Hepatol. 2018;69(1):43–50. PubMed PMID: 29454069. Epub 2018/02/18.

18. Gonzalez Martinez S, Molina Raya A, Becerra Massare A, et al. Liver transplantation in recipients with high model for end-stage liver disease score. Transplant Proc. 2018;50(2):595–7. PubMed PMID: 29579862. Epub 2018/03/28.

19. Cardoso FS, Karvellas CJ, Kneteman NM, et al. Postoperative resource utilization and survival among liver transplant recipients with model for end-stage liver disease score >/= 40: a retrospective cohort study. Can J Gastroenterol Hepatol. 2015;29(4):185–91. PubMed PMID: 25965438. Pubmed Central PMCID: PMC4444027. Epub 2015/05/13.

20. Petrowsky H, Rana A, Kaldas FM, et al. Liver transplantation in highest acuity recipients: identifying factors to avoid futility. Ann Surg. 2014;259(6):1186–94. PubMed PMID: 24263317. Epub 2013/11/23.

21. Panchal HJ, Durinka JB, Patterson J, et al. Survival outcomes in liver transplant recipients with model for end-stage liver disease scores of 40 or higher: a decade-long experience. HPB (Oxford). 2015;17(12):1074–84. PubMed PMID: 26373873. Pubmed Central PMCID: PMC4644359. Epub 2015/09/17.

22. Stratigopoulos P, Paul A, Hoyer DP, et al. High MELD score and extended operating time predict prolonged initial ICU stay after liver transplantation and influence the outcome. PLoS One. 2017;12(3):e0174173. PubMed PMID: 28319169. Pubmed Central PMCID: PMC5358862. Epub 2017/03/21.

23. Michard B, Artzner T, Lebas B, et al. Liver transplantation in critically ill patients: preoperative predictive factors of post-transplant mortality to avoid futility. Clin Transpl. 2017;31(12). PubMed PMID: 28895204. Epub 2017/09/13.

24. Nekrasov V, Matsuoka L, Kaur N, et al. Improvement in the outcomes of MELD >/= 40 liver transplantation: an analysis of 207 consecutive transplants in a highly competitive DSA. Transplantation. 2017;101(10):2360–7. PubMed PMID: 28319564. Epub 2017/03/21.

25. Nekrasov V, Matsuoka L, Rauf M, et al. National outcomes of liver transplantation for model for end-stage liver disease score >/=40: the impact of share 35. Am J Transplant. 2016;16(10):2912–24. PubMed PMID: 27063579. Epub 2016/04/12.
26. Volk ML, Lok AS, Pelletier SJ, et al. Impact of the model for end-stage liver disease allocation policy on the use of high-risk organs for liver transplantation. Gastroenterology. 2008;135(5):1568–74. PubMed PMID: 19009713. Epub 2008/11/15.
27. Bahra M, Neuhaus P. Liver transplantation in the high MELD era: a fair chance for everyone? Langenbeck's Arch Surg. 2011;396(4):461–5. PubMed PMID: 21384189. Epub 2011/03/09.
28. Bonney GK, Aldersley MA, Asthana S, et al. Donor risk index and MELD interactions in predicting long-term graft survival: a single-centre experience. Transplantation. 2009;87(12):1858–63. PubMed PMID: 19543065. Epub 2009/06/23.
29. Goldberg DS, Levine M, Karp S, et al. Share 35 changes in center-level liver acceptance practices. Liver Transpl. 2017;23(5):604–13. PubMed PMID: 28240804. Pubmed Central PMCID: PMC5462450. Epub 2017/02/28.
30. Brody BA, Haley A. Is futility a futile concept? J Med Philos. 1995;20(2):123–44. https://doi.org/10.1093/jmp/20.2.123.
31. Tomlinson T, Brody H. Ethics and communication in do-not-resuscitate orders. N Engl J Med. 1988;318(1):43–6. PubMed PMID: 3336383. Epub 1988/01/07.
32. Grant SB, Modi PK, Singer EA. Futility and the care of surgical patients: ethical dilemmas. World J Surg. 2014;38(7):1631–7. PubMed PMID: 24849199. Pubmed Central PMCID: PMC5176346. Epub 2014/05/23.
33. Schneiderman LJ, Jecker NS, Jonsen AR. Medical futility: its meaning and ethical implications. Ann Intern Med. 1990;112(12):949–54. PubMed PMID: 2187394. Epub 1990/06/15.
34. Rana A, Hardy MA, Halazun KJ, et al. Survival outcomes following liver transplantation (SOFT) score: a novel method to predict patient survival following liver transplantation. Am J Transplant. 2008;8(12):2537–46. PubMed PMID: 18945283. Epub 2008/10/24.
35. Linecker M, Krones T, Berg T, et al. Potentially inappropriate liver transplantation in the era of the “sickest first” policy – a search for the upper limits. J Hepatol. 2018;68(4):798–813. https://doi.org/10.1016/j.jhep.2017.11.008. Epub 2017 Nov 11.
36. Krowka MJ. Hepatopulmonary syndrome versus portopulmonary hypertension: distinctions and dilemmas. Hepatology. 1997;25(5):1282–4. PubMed PMID: 9141454. Epub 1997/05/01.
37. Yang YY, Lin HC, Lee WC, et al. Portopulmonary hypertension: distinctive hemodynamic and clinical manifestations. J Gastroenterol. 2001;36(3):181–6. PubMed PMID: 11291881. Epub 2001/04/09.
38. Salgia RJ, Goodrich NP, Simpson H, et al. Outcomes of liver transplantation for porto-pulmonary hypertension in model for end-stage liver disease era. Dig Dis Sci. 2014;59(8):1976–82. PubMed PMID: 24557576. Pubmed Central PMCID: PMC4119507. Epub 2014/02/22.
39. Rodriguez-Roisin R, Krowka MJ, Herve P, et al. Pulmonary-hepatic vascular disorders (PHD). Eur Respir J. 2004;24(5):861–80. PubMed PMID: 15516683. Epub 2004/11/02.
40. Krowka MJ, Frantz RP, McGoon MD, et al. Improvement in pulmonary hemodynamics during intravenous epoprostenol (prostacyclin): a study of 15 patients with moderate to severe portopulmonary hypertension. Hepatology. 1999;30(3):641–8. PubMed PMID: 10462369. Epub 1999/08/26.
41. Krowka MJ, Plevak DJ, Findlay JY, et al. Pulmonary hemodynamics and perioperative cardiopulmonary-related mortality in patients with portopulmonary hypertension undergoing liver transplantation. Liver Transpl. 2000;6(4):443–50. PubMed PMID: 10915166. Epub 2000/07/29.
42. Krowka MJ, Miller DP, Barst RJ, et al. Portopulmonary hypertension: a report from the US-based REVEAL Registry. Chest. 2012;141(4):906–15. PubMed PMID: 21778257. Epub 2011/07/23.
43. Swanson KL, Wiesner RH, Nyberg SL, et al. Survival in portopulmonary hypertension: Mayo Clinic experience categorized by treatment subgroups. Am J Transplant. 2008;8(11):2445–53. PubMed PMID: 18782292. Epub 2008/09/11.
44. Freeman RB Jr, Gish RG, Harper A, et al. Model for end-stage liver disease (MELD) exception guidelines: results and recommendations from the MELD Exception Study Group and Conference (MESSAGE) for the approval of patients who need liver transplantation with
diseases not considered by the standard MELD formula. Liver Transpl. 2006;12(12 Suppl 3):S128–36. PubMed PMID: 17123284. Epub 2006/11/24.

45. Krowka MJ, Wiesner RH, Heimbach JK. Pulmonary contraindications, indications and MELD exceptions for liver transplantation: a contemporary view and look forward. J Hepatol. 2013;59(2):367–74. PubMed PMID: 23557870. Epub 2013/04/06.

46. DuBrock HM, Goldberg DS, Sussman NL, et al. Predictors of waitlist mortality in portopulmonary hypertension. Transplantation. 2017;101(7):1609–15. PubMed PMID: 28207639. Pubmed Central PMCID: PMC5481480. Epub 2017/02/17.

47. Rajaram P, Parekh A, Fisher M, et al. Comparison of post-liver transplantation outcomes in portopulmonary hypertenion and pulmonary venous hypertension: a single-center experience. Transplant Proc. 2017;49(2):338–43. PubMed PMID: 28219595. Epub 2017/02/22.

48. Huang B, Shi Y, Liu J, et al. The early outcomes of candidates with portopulmonary hypertension after liver transplantation. BMC Gastroenterol. 2018;18(1):79. PubMed PMID: 29879915. Pubmed Central PMCID: PMC5992875. Epub 2018/06/09.

49. Sithamparanathan S, Nair A, Thirugnanasothy L, et al. Survival in portopulmonary hypertension: outcomes of the United Kingdom National Pulmonary Arterial Hypertension Registry. J Heart Lung Transplant. 2017;36(7):770–9. PubMed PMID: 28190786. Epub 2017/02/14.

50. Nadim MK, Sung RS, Davis CL, et al. Simultaneous liver-kidney transplantation summit: current state and future directions. Am J Transplant. 2012;12(11):2901–8. PubMed PMID: 22822723. Epub 2012/07/25.

51. Eason JD, Gonwa TA, Davis CL, et al. Proceedings of consensus conference on simultaneous liver kidney transplantation (SLK). Am J Transplant. 2008;8(11):2243–51. PubMed PMID: 18808402. Epub 2008/09/24.

52. Formica RN, Aeder M, Boyle G, et al. Simultaneous liver-kidney allocation policy: a proposal to optimize appropriate utilization of scarce resources. Am J Transplant. 2016;16(3):758–66. PubMed PMID: 26603142. Epub 2015/11/26.

53. Lunsford KE, Bodzin AS, Markovic D, et al. Avoiding futility in simultaneous liver-kidney transplantation: analysis of 331 consecutive patients listed for dual organ replacement. Ann Surg. 2017;265(5):1016–24. PubMed PMID: 27232249. Epub 2016/05/28.

54. Fong TL, Bunnapradist S, Jordan SC, et al. Analysis of the united network for organ sharing database comparing renal allografts and patient survival in combined liver-kidney transplantation with the contralateral allografts in kidney alone or kidney-pancreas transplantation. Transplantation. 2003;76(2):348–53. PubMed PMID: 12883191. Epub 2003/07/29.

55. Simpson N, Cho YW, Cicciarelli JC, et al. Comparison of renal allograft outcomes in combined liver-kidney transplantation versus subsequent kidney transplantation in liver transplant recipients: analysis of UNOS database. Transplantation. 2006;82(10):1298–303. PubMed PMID: 17130778. Epub 2006/11/30.

56. Puri V, Eason J. Simultaneous liver-kidney transplantation. Curr Transplant Rep. 2015;2(4):297–302. PubMed PMID: 26523249. Pubmed Central PMCID: PMC4623067. Epub 2015/11/03.

57. Davis CL, Feng S, Sung R, et al. Simultaneous liver-kidney transplantation: evaluation to decision making. Am J Transplant. 2007;7(7):1702–9. PubMed PMID: 17532752. Epub 2007/05/30.

58. Ceulemans LJ, Strypstein S, Neyrinck A, et al. Combined liver-thoracic transplantation: single-center experience with introduction of the 'Liver-first' principle. Transpl Int. 2016;29(6):715–26. PubMed PMID: 27037837. Epub 2016/04/03.

59. Beal EW, Mumtaz K, Hayes D Jr, et al. Combined heart-liver transplantation: indications, outcomes and current experience. Transplant Rev (Orlando). 2016;30(4):261–8. PubMed PMID: 27527917. Pubmed Central PMCID: PMC5450025. Epub 2016/08/17.

60. Cannon RM, Hughes MG, Jones CM, et al. A review of the United States experience with combined heart-liver transplantation. Transpl Int. 2012;25(12):1223–8. PubMed PMID: 22937819. Epub 2012/09/04.
61. Careddu L, Zanfi C, Pantaleo A, et al. Combined heart-liver transplantation: a single-center experience. Transpl Int. 2015;28(7):828–34. PubMed PMID: 25711771. Epub 2015/02/26.
62. Wong TW, Gandhi MJ, Daly RC, et al. Liver allograft provides Immunoprotection for the cardiac allograft in combined heart-liver transplantation. Am J Transplant. 2016;16(12):3522–31. PubMed PMID: 27184686. Epub 2016/05/18.
63. Weill D. Lung transplantation: indications and contraindications. J Thorac Dis. 2018;10(7):4574–87. PubMed PMID: 30174910. Pubmed Central PMCID: PMC6105990. Epub 2018/09/04.
64. Gadre S, Turowski J, Budev M. Overview of lung transplantation, heart-lung transplantation, and combined hematopoietic stem cell transplantation and lung transplantation. Clin Chest Med. 2017;38(4):623–40. PubMed PMID: 29128014. Epub 2017/11/13.
65. Pinderski LJ, Kirklin JK, McGiffin D, et al. Multi-organ transplantation: is there a protective effect against acute and chronic rejection? J Heart Lung Transplant. 2005;24(11):1828–33. PubMed PMID: 16297789. Epub 2005/11/22.
66. Rana A, Robles S, Russo MJ, et al. The combined organ effect: protection against rejection? Ann Surg. 2008;248(5):871–9. PubMed PMID: 18948817. Epub 2008/10/25.
67. Gottlieb J. Lung allocation. J Thorac Dis. 2017;9(8):2670–4. PubMed PMID: 28932574. Pubmed Central PMCID: PMC564149. Epub 2017/09/22.
68. Wolf JH, Sulewski ME, Cassuto JR, et al. Simultaneous thoracic and abdominal transplantation: can we justify two organs for one recipient? Am J Transplant. 2013;13(7):1806–16. PubMed PMID: 23718142. Epub 2013/05/31.
69. Grannas G, Neipp M, Hoepf MM, et al. Indications for and outcomes after combined lung and liver transplantation: a single-center experience on 13 consecutive cases. Transplantation. 2008;85(4):524–31. PubMed PMID: 18347530. Epub 2008/03/19.
70. Yi SG, Burroughs SG, Loebe M, et al. Combined lung and liver transplantation: analysis of a single-center experience. Liver Transpl. 2014;20(1):46–53. PubMed PMID: 24136814. Epub 2013/10/19.
71. Scheiermann P, Czerner S, Kaspar M, et al. Combined lung and liver transplantation with extracorporeal membrane oxygenation instead of cardiopulmonary bypass. J Cardiothorac Vasc Anesth. 2016;30(2):437–42. PubMed PMID: 26432697. Epub 2015/10/04.
72. Butler P, Israel L, Nusbacher J, et al. Blood transfusion in liver transplantation. Transfusion. 1985;25(2):120–3. PubMed PMID: 3885484. Pubmed Central PMCID: PMC2967283. Epub 1985/03/01.
73. Massicotte L, Denault AY, Beaulieu D, et al. Transfusion rate for 500 consecutive liver transplantations: experience of one liver transplantation center. Transplantation. 2012;93(12):1276–81. PubMed PMID: 22617090. Epub 2012/05/24.
74. Schroeder RA, Kuo PC. Pro: low central venous pressure during liver transplantation – not too low. J Cardiothorac Vasc Anesth. 2008;22(2):311–4. PubMed PMID: 18375341. Epub 2008/04/01.
75. Massicotte L, Lenis S, Thibeault L, et al. Reduction of blood product transfusions during liver transplantation. Can J Anaesth. 2005;52(5):545–6. PubMed PMID: 15872137. Epub 2005/05/06.
76. Xia VW, Fond A, Du B. Ascites, but not hyponatremia, is associated with high intraoperative transfusion and vasopressor requirements during liver transplantation. Transplant Proc. 2006;38(5):1398–9. PubMed PMID: 16797315. Epub 2006/06/27.
77. Esmat Gamal M, Pirene J, Van Malenestien H, et al. Risk factors for bleeding and clinical implications in patients undergoing liver transplantation. Transplant Proc. 2012;44(9):2857–60. PubMed PMID: 23146542. Epub 2012/11/14.
78. Massicotte L, Sasseine MP, Lenis S, et al. Survival rate changes with transfusion of blood products during liver transplantation. Can J Anaesth. 2005;52(2):148–55. PubMed PMID: 15684254. Epub 2005/02/03.
79. Ramos E, Dalmau A, Sabate A, et al. Intraoperative red blood cell transfusion in liver transplantation: influence on patient outcome, prediction of requirements, and measures to reduce them. Liver Transpl. 2003;9(12):1320–7. PubMed PMID: 14625833. Epub 2003/11/20.
80. Boin IF, Leonardi MI, Luzo AC, et al. Intraoperative massive transfusion decreases survival after liver transplantation. Transplant Proc. 2008;40(3):789–91. PubMed PMID: 18455018. Epub 2008/05/06.

81. Massicotte L, Sassi MP, Lenis S, Roy A. Transfusion predictors in liver transplant. Anesth Analg. 2004;98(5):1245–51, table of contents. PubMed PMID: 15105195. Epub 2004/04/24.

82. McCluskey SA, Karkouti K, Wijeysundera DN, et al. Derivation of a risk index for the prediction of massive blood transfusion in liver transplantation. Liver Transpl. 2006;12(11):1584–93. PubMed PMID: 16952177. Epub 2006/09/05.

83. Saracoglu A, Saracoglu KT. Coagulopathy during liver transplantation. J Anaesthesiol Clin Pharmacol. 2018;34(3):289–95. PubMed PMID: 30386008. Pubmed Central PMCID: PMC6194832. Epub 2018/11/06.

84. Violi F, Basili S, Raparelli V, et al. Patients with liver cirrhosis suffer from primary haemostatic defects? Fact or fiction? J Hepatol. 2011;55(6):1415–27. PubMed PMID: 21718668. Epub 2011/07/02.

85. Lisman T, Porte RJ. Rebalanced hemostasis in patients with liver disease: evidence and clinical consequences. Blood. 2010;116(6):878–85. PubMed PMID: 20400681. Epub 2010/04/20.

86. Tripodi A, Salerno F, Chantarangkul V, et al. Evidence of normal thrombin generation in cirrhosis despite abnormal conventional coagulation tests. Hepatology. 2005;41(3):553–8. PubMed PMID: 15726661. Epub 2005/02/24.

87. Caldwell SH, Hoffman M, Lisman T, et al. Coagulation disorders and hemostasis in liver disease: pathophysiology and critical assessment of current management. Hepatology. 2006;44(4):1039–46. PubMed PMID: 17006940. Epub 2006/09/29.

88. Cleland S, Corredor C, Ye JJ, et al. Massive haemorrhage in liver transplantation: consequences, prediction and management. World J Transplant. 2016;6(2):291–305. PubMed PMID: 27358774. Pubmed Central PMCID: PMC4919733. Epub 2016/07/01.

89. Calne RY. Surgical aspects of clinical liver transplantation in 14 cases. Br J Surg. 1969;56(10):729–36. PubMed PMID: 4899842. Epub 1969/10/01.

90. Steib A, Freys G, Lehmnn C, et al. Intraoperative blood losses and transfusion requirements during adult liver transplantation remain difficult to predict. Can J Anaesth. 2001;48(11):1075–9. PubMed PMID: 11744582. Epub 2001/12/18.

91. Reyle-Hahn M, Rossaint R. Coagulation techniques are not important in directing blood product transfusion during liver transplantation. Liver Transpl Surg. 1997;3(6):659–63; discussion 63–5. PubMed PMID: 9404973. Epub 1997/12/24.

92. Adelmann D, Kronish K, Ramsay MA. Anesthesia for liver transplantation. Anesthesiol Clin. 2017;35(3):491–508. PubMed PMID: 28784222. Epub 2017/08/09.

93. Lisman T, Porte RJ. Platelet function in patients with cirrhosis. J Hepatol. 2012;56(4):993–4; author reply 4–5. PubMed PMID: 22424439. Epub 2012/03/20.

94. Donohue CI, Mallett SV. Reducing transfusion requirements in liver transplantation. World J Transplant. 2015;5(4):165–82. PubMed PMID: 26722645. Pubmed Central PMCID: PMC4689928. Epub 2016/01/02.

95. Tzakis A, Todo S, Starzl TE. Orthotopic liver transplantation with preservation of the inferior vena cava. Ann Surg. 1989;210(5):649–52. PubMed PMID: 2818033. Pubmed Central PMCID: PMC1357802. Epub 1989/11/01.

96. Belghiti J, Panis Y, Sauvanet A, et al. A new technique of side to side caval anastomosis during orthotopic hepatic transplantation without inferior vena caval occlusion. Surg Gynecol Obstet. 1992;175(3):270–2. PubMed PMID: 1514163. Epub 1992/09/01.

97. Schmitz V, Schoening W, Jelkmann I, et al. Different cava reconstruction techniques in liver transplantation: piggyback versus cava resection. Heparntobilary Pancreat Dis Int. 2014;13(3):242–9. PubMed PMID: 24919606. Epub 2014/06/13.

98. Moreno-Gonzalez E, Menene-Diaz JG, Fendura Y, et al. Advantages of the piggy back technique on intraoperative transfusion, fluid compsumption, and vasoactive drugs requirements in liver transplantation: a comparative study. Transplant Proc. 2003;35(5):1918–9. PubMed PMID: 12962848. Epub 2003/09/10.
99. Navarro F, Le Moine MC, Fabre JM, et al. Specific vascular complications of orthotopic liver transplantation with preservation of the retrohepatic vena cava: review of 1361 cases. Transplantation. 1999;68(5):646–50. PubMed PMID: 10507483. Epub 1999/10/03.

100. Forkin KT, Colquhoun DA, Nemergut EC, Huffmyer JL. The coagulation profile of end-stage liver disease and considerations for intraoperative management. Anesth Analg. 2018;126(1):46–61. PubMed PMID: 28795966. Epub 2017/08/11.

101. Thakrar SV, Melikian CN. Anaesthesia for liver transplantation. Br J Hosp Med (Lond). 2017;78(5):260–5. PubMed PMID: 28489450. Epub 2017/05/11.

102. Hilmi I, Horton CN, Planinsic RM, et al. The impact of postreperfusion syndrome on short-term patient and liver allograft outcome in patients undergoing orthotopic liver transplantation. Liver Transpl. 2008;14(4):504–8. PubMed PMID: 18383079. Epub 2008/04/03.

103. Siniscalchi A, Gamberini L, Laici C, et al. Post reperfusion syndrome during liver transplantation: from pathophysiology to therapy and preventive strategies. World J Gastroenterol. 2016;22(4):1551–69. PubMed PMID: 26819522. Pubmed Central PMCID: PMC4721988. Epub 2016/01/29.

104. Brems JJ, Millis JM, Hiatt JR, et al. Hepatic artery reconstruction during liver transplantation. Transplantation. 1989;47(2):403–6. PubMed PMID: 2645728. Epub 1989/02/01.

105. Mallett SV. Clinical utility of viscoelastic tests of coagulation (TEG/ROTEM) in patients with liver disease and during liver transplantation. Semin Thromb Hemost. 2015;41(5):527–37. PubMed PMID: 26049072. Epub 2015/06/07.

106. Hawkins RB, Raymond SL, Hartjes T, et al. Review: the perioperative use of thromboelastography for liver transplant patients. Transplant Proc. 2018;50(10):3552–8. PubMed PMID: 30577236. Epub 2018/12/24.

107. Ganter MT, Hofer CK. Coagulation monitoring: current techniques and clinical use of viscoelastic point-of-care coagulation devices. Anesth Analg. 2008;106(5):1366–75. PubMed PMID: 18420846. Epub 2008/04/19.

108. Lawson PJ, Moore HB, Moore EE, et al. Preoperative thrombelastography maximum amplitude predicts massive transfusion in liver transplantation. J Surg Res. 2017;220:171–5. PubMed PMID: 29180179. Pubmed Central PMCID: PMC5726438. Epub 2017/11/29.

109. Tafur LA, Taura P, Blasi A, et al. Rotation thromboelastometry velocity curve predicts blood loss during liver transplantation. Br J Anaesth. 2016;117(6):741–8. PubMed PMID: 27956672. Epub 2016/12/14.

110. Wang SC, Shieh JF, Chang KY, et al. Thromboelastography-guided transfusion decreases intraoperative blood transfusion during orthotopic liver transplantation: randomized clinical trial. Transplant Proc. 2010;42(7):2590–3. PubMed PMID: 20832550. Epub 2010/09/14.

111. Zamper RPC, Amorim TC, Queiroz VNF, et al. Association between viscoelastic tests-guided therapy with synthetic factor concentrates and allogenic blood transfusion in liver transplantation: a before-after study. BMC Anesthesiol. 2018;18(1):198. PubMed PMID: 30579327. Pubmed Central PMCID: PMC6303918. Epub 2018/12/24.

112. Molenaar IQ, Warnaar N, Groen H, et al. Efficacy and safety of antifibrinolytic drugs in liver transplantation: a systematic review and meta-analysis. Am J Transplant. 2007;7(1):185–94. PubMed PMID: 17227567. Epub 2007/01/18.

113. Kozek-Langenecker SA, Afshari A, Albaladejo P, et al. Management of severe perioperative bleeding: guidelines from the European Society of Anaesthesiology. Eur J Anaesthesiol. 2013;30(6):270–382. PubMed PMID: 23656742. Epub 2013/05/10.

114. Trzebicki J, Flakiewicz E, Kosieradzki M, et al. The use of thromboelastometry in the assessment of hemostasis during orthotopic liver transplantation reduces the demand for blood products. Ann Transplant. 2010;15(3):19–24. PubMed PMID: 20877262. Epub 2010/09/30.

115. Henderson JM, Mackay GJ, Hooks M, et al. High cardiac output of advanced liver disease persists after orthotopic liver transplantation. Hepatology. 1992;15(2):258–62. PubMed PMID: 1735528. Epub 1992/02/01.

116. Hori T, Ogura Y, Onishi Y, et al. Systemic hemodynamics in advanced cirrhosis: concerns during perioperative period of liver transplantation. World J Hepatol. 2016;8(25):1047–60. PubMed PMID: 27660671. Pubmed Central PMCID: PMC5026996. Epub 2016/09/24.
117. Piardi T, Lhuaire M, Bruno O, et al. Vascular complications following liver transplantation: a literature review of advances in 2015. World J Hepatol. 2016;8(1):36–57. PubMed PMID: 26783420. Pubmed Central PMCID: PMC4705452. Epub 2016/01/20.

118. Heaton ND. Hepatic artery thrombosis: conservative management or retransplantation? Liver Transpl. 2013;19(Suppl 2):S14–6. PubMed PMID: 24019107. Epub 2013/09/11.

119. Singhal A, Stokes K, Sebastian A, et al. Endovascular treatment of hepatic artery thrombosis following liver transplantation. Transpl Int. 2010;23(3):245–56. PubMed PMID: 20030796. Epub 2009/12/25.

120. Buchholz BM, Khan S, David MD, et al. Retransplantation in late hepatic artery thrombosis: graft access and transplant outcome. Transplant Direct. 2017;3(8):e186. PubMed PMID: 28795138. Pubmed Central PMCID: PMC5540624. Epub 2017/08/11.

121. Bekker J, Ploem S, de Jong KP. Early hepatic artery thrombosis after liver transplantation: a systematic review of the incidence, outcome and risk factors. Am J Transplant. 2009;9(4):746–57. PubMed PMID: 19298450. Epub 2009/03/21.

122. Gunsar F, Rolando N, Pastacaldi S, et al. Late hepatic artery thrombosis after orthotopic liver transplantation. Liver Transpl. 2010;16(6):605–11. PubMed PMID: 20030796. Epub 2009/12/25.

123. Dani G, Sun MR, Bennett AE. Imaging of liver transplant and its complications. Semin Ultrasound CT MR. 2013;34(4):365–77. PubMed PMID: 23895908. Epub 2013/07/31.

124. Herrero A, Souche R, Joly E, et al. Early hepatic artery thrombosis after liver transplantation: what is the impact of the arterial reconstruction type? World J Surg. 2017;41(8):2101–10. PubMed PMID: 28324141. Epub 2017/03/23.

125. Chatzizacharias NA, Aly M, Praseedom RK. The role of arterial conduits for revascularisation in adult orthotopic liver transplantation. Transplant Rev (Orlando). 2017;31(2):121–6. PubMed PMID: 27884502. Epub 2016/11/26.

126. Schroering JR, Kubal CA, Fridell JA, et al. Impact of variant donor hepatic arterial anatomy on clinical graft outcomes in liver transplantation. Liver Transpl. 2018;24(10):1481–4. PubMed PMID: 30054968. Pubmed Central PMCID: PMC6298596. Epub 2018/07/29.

127. Sheiner PA, Varma CV, Guerrera JV, et al. Selective revascularization of hepatic artery thromboses after liver transplantation improves patient and graft survival. Transplantation. 1997;64(9):1295–9. PubMed PMID: 9371671. Epub 1997/11/26.

128. Crossin JD, Muradali D, Wilson SR. US of liver transplants: normal and abnormal. Radiographics. 2003;23(5):1093–114. PubMed PMID: 12975502. Epub 2003/09/17.

129. Baheti AD, Sanyal R, Heller MT, Bhargava P. Surgical techniques and imaging complications of liver transplant. Radiol Clin N Am. 2016;54(2):199–215. PubMed PMID: 26896220. Epub 2016/02/21.

130. Marin-Gomez LM, Bernal-Bellido C, Alamo-Martinez JM, et al. Intraoperative hepatic artery blood flow predicts early hepatic artery thrombosis after liver transplantation. Transplant Proc. 2012;44(7):2078–81. PubMed PMID: 22974916. Epub 2012/09/15.

131. Stange BJ, Glanemann M, Nuessler NC, et al. Hepatic artery thrombosis after adult liver transplantation. Liver Transpl. 2003;9(6):612–20. PubMed PMID: 12783404. Epub 2003/06/05.

132. Scarinci A, Sainz-Barriga M, Berrevoet F, et al. Early arterial revascularization after hepatic artery thrombosis may avoid graft loss and improve outcomes in adult liver transplantation. Transplant Proc. 2010;42(10):4403–8. PubMed PMID: 21168708. Epub 2010/12/21.

133. Chen J, Weinstein J, Black S, et al. Surgical and endovascular treatment of hepatic arterial complications following liver transplant. Clin Transpl. 2014;28(12):1305–12. PubMed PMID: 25091402. Epub 2014/08/06.

134. Rostambeigi N, Hunter D, Duval S, et al. Stent placement versus angioplasty for hepatic artery stenosis after liver transplant: a meta-analysis of case series. Eur Radiol. 2013;23(5):1323–34. PubMed PMID: 23239061. Epub 2012/12/15.

135. Pitchaimuthu M, Roll GR, Zia Z, et al. Long-term follow-up after endovascular treatment of hepatic venous outflow obstruction following liver transplantation. Transpl Int. 2016;29(10):1106–16. PubMed PMID: 27371935. Epub 2016/07/03.
136. Umehara M, Narumi S, Sugai M, et al. Hepatic venous outflow obstruction in living donor liver transplantation: balloon angioplasty or stent placement? Transplant Proc. 2012;44(3):769–71. PubMed PMID: 22483491. Epub 2012/04/10.

137. Arudchelvam J, Bartlett A, McCall J, et al. Hepatic venous outflow obstruction in piggyback liver transplantation: single centre experience. ANZ J Surg. 2017;87(3):182–5. PubMed PMID: 26471387. Epub 2015/10/17.

138. Khorsandi SE, Athale A, Vilca-Melendez H, et al. Presentation, diagnosis, and management of early hepatic venous outflow complications in whole cadaveric liver transplant. Liver Transpl. 2015;21(7):914–21. PubMed PMID: 25907399. Epub 2015/04/25.

139. Fujimori M, Yamakado K, Takaki H, et al. Long-term results of stent placement in patients with outflow block after living-donor-liver transplantation. Cardiovasc Intervent Radiol. 2016;39(4):566–74. PubMed PMID: 26464222. Epub 2015/10/16.

140. Wang SL, Sze DY, Busque S, et al. Treatment of hepatic venous outflow obstruction after piggyback liver transplantation. Radiology. 2005;236(1):352–9. PubMed PMID: 15955856. Epub 2005/06/16.

141. Kim KS, Lee JS, Choi GS, et al. Long-term outcomes after stent insertion in patients with early and late early vein outflow obstruction after living donor liver transplantation. Ann Surg Treat Res. 2018;95(6):333–9. PubMed PMID: 30505825. Pubmed Central PMCID: PMC6255746. Epub 2018/12/07.

142. Kochhar G, Parungao JM, Hanouneh IA, Parsi MA. Biliary complications following liver transplantation. World J Gastroenterol. 2013;19(19):2841–6. PubMed PMID: 23704818. Pubmed Central PMCID: PMC3660810. Epub 2013/05/25.

143. Axelrod DA, Dzebisashvilli N, Lentine KL, et al. National assessment of early biliary complications after liver transplantation: economic implications. Transplantation. 2014;98(11):1226–35. PubMed PMID: 25191226. Epub 2014/08/15.

144. O'Neill S, Roebuck A, Khoo E, et al. A meta-analysis and meta-regression of outcomes including biliary complications in donation after cardiac death liver transplantation. Transpl Int. 2014;27(11):1159–74. PubMed PMID: 25052036. Epub 2014/07/24.

145. Seeboer D, Euirch Y, Veltzke-Schlieker W, Neuhaus P. Biliary complications after liver transplantation: old problems and new challenges. Am J Transplant. 2013;13(2):253–65. PubMed PMID: 23331505. Epub 2013/01/22.

146. Kim WR, Lake JR, Smith JM, et al. OPTN/SRTR 2013 annual data report: liver. Am J Transplant. 2015;15(Suppl 2):1–28. PubMed PMID: 25623641. Epub 2015/01/30.

147. Dutkowski P, Oberkofer CE, Slankamenac K, et al. Are there better guidelines for allocation in liver transplantation? A novel score targeting justice and utility in the model for end-stage liver disease era. Ann Surg. 2011;254(5):745–53; discussion 53. PubMed PMID: 22042468. Epub 2011/11/02.

148. Feng S, Goodrich NP, Bragg-Gresham JL, et al. Characteristics associated with liver graft failure: the concept of a donor risk index. Am J Transplant. 2006;6(4):783–90. PubMed PMID: 16539636. Epub 2006/03/17.

149. Halldorson JB, Bakhvatsalam R, Fix O, et al. D-MELD, a simple predictor of post liver transplant mortality for optimization of donor/recipient matching. Am J Transplant. 2009;9(2):318–26. PubMed PMID: 19120079. Epub 2009/01/06.

150. Northup PG, Berg CL. Preoperative delta-MELD score does not independently predict mortality after liver transplantation. Am J Transplant. 2004;4(10):1643–9. PubMed PMID: 15367219. Epub 2004/09/16.

151. Briceno J, Ciria R, de la Mata M. Donor-recipient matching: myths and realities. J Hepatol. 2013;58(4):811–20. PubMed PMID: 23041646. Epub 2012/10/30.

152. Buescher N, Seeboer D, Helbig M, et al. Evaluating twenty-years of follow-up after orthotopic liver transplantation, best practice for donor-recipient matching: what can we learn from the past era? World J Transplant. 2016;6(3):599–607. PubMed PMID: 27683639. Pubmed Central PMCID: PMC5036130. Epub 2016/09/30.
153. Rauchfuss F, Zidan A, Scheuerlein H, et al. Waiting time, not donor-risk-index, is a major determinant for beneficial outcome after liver transplantation in high-MELD patients. Ann Transplant. 2013;18:243–7. PubMed PMID: 23792527. Epub 2013/06/26.

154. Englesbe MJ, Kubus J, Muhammad W, et al. Portal vein thrombosis and survival in patients with cirrhosis. Liver Transpl. 2010;16(1):83–90. PubMed PMID: 20035521. Epub 2009/12/26.

155. Paskonis M, Jurgaitis J, Mehrabi A, et al. Surgical strategies for liver transplantation in the case of portal vein thrombosis--current role of cavoportal hemitransposition and renportal anastomosis. Clin Transpl. 2006;20(5):551–62. PubMed PMID: 16968480. Epub 2006/09/14.

156. Ögren M, Bergqvist D, Björck M et al. Portal vein thrombosis: prevalence, patient characteristics and lifetime risk: a population study based on 23,796 consecutive autopsies. World J Gastroenterol. 2006 Apr 7;12(13):2115–9. https://doi.org/10.3748/wjg.v12.i13.2115.

157. Ögren M, Bergqvist D, Björck M, et al. Portal vein thrombosis: prevalence, patient characteristics and lifetime risk: a population study based on 23,796 consecutive autopsies. World J Gastroenterol. 2006;12(13):2115–9. https://doi.org/10.3748/wjg.v12.i13.2115.

158. Amitrano L, Brancaccio V, Guardascione MA, et al. Inherited coagulation disorders in cirrhotic patients with portal vein thrombosis. Hepatology. 2000;31(2):345–8. PubMed PMID: 10655256. Epub 2000/02/03.

159. Chamarthy MR, Anderson ME, Pillai AK, Kalva SP. Thrombolysis and Transjugular intrahepatic Portosystemic shunt creation for acute and subacute portal vein thrombosis. Tech Vasc Interv Radiol. 2016;19(1):42–51. PubMed PMID: 26997088. Epub 2016/03/22.

160. Ponziani FR. Portal vein thrombosis: insight into physiopathology, diagnosis, and treatment. World J Gastroenterol. 2010;16(2):143–55.

161. Valla DC, Condat B, Lobrec D. Spectrum of portal vein thrombosis in the west. J Gastroenterol Hepatol. 2002;17(Suppl 3):S224–7. PubMed PMID: 12472940. Epub 2002/12/11.

162. Chawla Y, Duseja A, Dhiman RK. Review article: the modern management of portal vein thrombosis. Aliment Pharmacol Ther. 2009;30(9):881–94. PubMed PMID: 19678814. Epub 2009/08/15.

163. Delgado MG, Seijo S, Yepes I, et al. Efficacy and safety of anticoagulation on patients with cirrhosis and portal vein thrombosis. Clin Gastroenterol Hepatol. 2012;10(7):776–83. PubMed PMID: 22289875. Epub 2012/02/01.

164. Villa E, Camma C, Marietta M, et al. Enoxaparin prevents portal vein thrombosis and liver deccompensation in patients with advanced cirrhosis. Gastroenterology. 2012;143(5):1253–60 e4. PubMed PMID: 22819864. Epub 2012/07/24.

165. Thornburg B, Desai K, Hickey R, et al. Portal vein recanalization and transjugular intrahepatic portosystemic shunt creation for chronic portal vein thrombosis: technical considerations. Tech Vasc Interv Radiol. 2016;19(1):52–60. PubMed PMID: 26997089. Epub 2016/03/22.

166. Luca A, Miraglia R, Caruso S, et al. Short- and long-term effects of the transjugular intrahepatic portosystemic shunt on portal vein thrombosis in patients with cirrhosis. Gut. 2011;60(6):846–52. PubMed PMID: 21357252. Epub 2011/03/02.

167. Manzanet G, Sanjuan F, Orbis P, et al. Liver transplantation in patients with portal vein thrombosis. Liver Transpl. 2001;7(2):125–31. PubMed PMID: 11172396. Epub 2001/02/15.

168. Shaked A, Busuttil RW. Liver transplantation in patients with portal vein thrombosis and central portacaval shunts. Ann Surg. 1991;214(6):696–702. PubMed PMID: 1741649. Pubmed Central PMCID: PMC1358494. Epub 1991/12/01.

169. Englesbe MJ, Schaubel DE, Cai S, et al. Portal vein thrombosis and liver transplant survival benefit. Liver Transpl. 2010;16(8):999–1005. PubMed PMID: 20677291. Pubmed Central PMCID: PMC2915450. Epub 2010/08/03.

170. Doenecke A, Tsui TY, Zuelke C, et al. Pre-existent portal vein thrombosis in liver transplantation: influence of pre-operative disease severity. Clin Transpl. 2010;24(1):48–55. PubMed PMID: 19236435. Epub 2009/02/25.

171. Molmenti EP, Roodhouse TW, Molmenti H, et al. Thrombendarcovenectomy for organized portal vein thrombosis at the time of liver transplantation. Ann Surg. 2002;235(2):292–6. PubMed PMID: 11807371. Pubmed Central PMCID: PMC1422428. Epub 2002/01/25.
172. Gimeno FA, Calvo J, Loinaz C, et al. Comparative analysis of the results of orthotopic liver transplantation in patients with and without portal vein thrombosis. Transplant Proc. 2005;37(9):3899–903. PubMed PMID: 16386578. Epub 2006/01/03.

173. Llado L, Fabregat J, Castellote J, et al. Management of portal vein thrombosis in liver transplantation: influence on morbidity and mortality. Clin Transpl. 2007;21(6):716–21. PubMed PMID: 17988264. Epub 2007/11/09.

174. Yerdel MA, Gunson B, Mirza D, et al. Portal vein thrombosis in adults undergoing liver transplantation: risk factors, screening, management, and outcome. Transplantation. 2000;69(9):1873–81. PubMed PMID: 10830225. Epub 2000/06/01.

175. Pan C, Shi Y, Zhang JJ, et al. Single-center experience of 253 portal vein thrombosis patients undergoing liver transplantation in China. Transplant Proc. 2009;41(9):3761–5. PubMed PMID: 19917382. Epub 2009/11/18.

176. Jia YP, Lu Q, Gong S, et al. Postoperative complications in patients with portal vein thrombosis after liver transplantation: evaluation with Doppler ultrasonography. World J Gastroenterol. 2007;13(34):4636–40. PubMed PMID: 17729421. Pubmed Central PMCID: PMC4611842. Epub 2007/08/31.

177. Quintini C, Spaggiari M, Hashimoto K, et al. Safety and effectiveness of renoportal bypass in patients with complete portal vein thrombosis: an analysis of 10 patients. Liver Transpl. 2015;21(3):344–52. PubMed PMID: 25420619. Epub 2014/11/26.

178. Hibi T, Nishida S, Levi DM, et al. When and why portal vein thrombosis matters in liver transplantation: a critical audit of 174 cases. Ann Surg. 2014;259(4):760–6. PubMed PMID: 24299686. Epub 2013/12/05.

179. Lautt WW. Regulatory processes interacting to maintain hepatic blood flow constancy: vascular compliance, hepatic arterial buffer response, hepatorenal reflex, liver regeneration, escape from vasoconstriction. Hepatol Res. 2007;37(11):891–903. PubMed PMID: 17854463. Pubmed Central PMCID: PMC2981600. Epub 2007/09/15.

180. Funaki B, Rosenblum JD, Leef JA, et al. Portal vein stenosis in children with segmental liver transplants: treatment with percutaneous transhepatic venoplasty. AJR Am J Roentgenol. 1995;165(1):161–5. PubMed PMID: 7785578. Epub 1995/07/01.

181. Shibata T, Itoh K, Kubo T, et al. Percutaneous transhepatic balloon dilation of portal venous stenosis in patients with living donor liver transplantation. Radiology. 2005;235(3):1078–83. PubMed PMID: 15845790. Epub 2005/04/23.

182. Lautt WW. The 1995 Ciba-Geigy Award Lecture. Intrinsic regulation of hepatic blood flow. Can J Physiol Pharmacol. 1996;74(3):223–33. PubMed PMID: 8773400. Epub 1996/03/01.

183. Jin WX, Wang B, Zhang YL, et al. Effects of hepatic blood inflow on liver ultrastructure and regeneration after extensive liver resection in rats with cirrhosis. Exp Ther Med. 2018;16(3):2573–83. PubMed PMID: 30210605. Pubmed Central PMCID: PMC6122590. Epub 2018/09/14.

184. Kim PTW, Klintmalm GB. J Hepatol Gastroint Dis 2016, 2:2 https://doi.org/10.4172/jhgd.1000127.

185. Kim PTW, Klintmalm GB. J Hepatol Gastroint Dis. 2016;2(2) https://doi.org/10.4172/jhgd.1000127.
190. Smyrniotis V, Kostopanagiotou G, Kondi A, et al. Hemodynamic interaction between portal vein and hepatic artery flow in small-for-size split liver transplantation. Transpl Int. 2002;15(7):355–60. PubMed PMID: 12122512. Epub 2002/07/18.

191. Aoki T, Imamura H, Kaneko J, et al. Intraoperative direct measurement of hepatic arterial buffer response in patients with or without cirrhosis. Liver Transpl. 2005;11(6):684–91. PubMed PMID: 15915492. Epub 2005/05/26.

192. Akamatsu N, Sugawara Y, Satou S, et al. Hemodynamic changes in the hepatic circulation after the modulation of the splenic circulation in an in vivo human experimental model. Liver Transpl. 2014;20(1):116–21. PubMed PMID: 24123877. Epub 2013/10/15.

193. Saad WE. Nonocclusive hepatic artery hypoperfusion syndrome (splenic steal syndrome) in liver transplant recipients. Semin Intervent Radiol. 2012;29(2):140–6. PubMed PMID: 23729985. Pubmed Central PMCID: PMC3444879. Epub 2013/06/05.

194. Iranpour P, Lall C, Houshyar R, et al. Altered Doppler flow patterns in cirrhosis patients: an overview. Ultrasonography. 2016;35(1):3–12. PubMed PMID: 26169079. Pubmed Central PMCID: PMC4701371. Epub 2015/07/15.

195. Chang PD, Rajeswaran S, Nikolaidis P, et al. Cirrhotic right and left portal veins: how slow do they go? Identification of threshold velocities associated with subsequent development of hepatofugal flow. Ultrasound Q. 2013;29(2):131–5. PubMed PMID: 23698619. Epub 2013/05/24.

196. Kelly DM, Zhu X, Shibah, et al. Adenosine restores the hepatic artery buffer response and improves survival in a porcine model of small-for-size syndrome. Liver Transpl. 2009;15(11):1448–57. PubMed PMID: 19877203. Epub 2009/10/31.

197. Bolognesi M, Sacerdoti D, Bombonato G, et al. Change in portal flow after liver transplantation: effect on hepatic arterial resistance indices and role of spleen size. Hepatology. 2002;35(3):601–8. PubMed PMID: 11870373. Epub 2002/03/01.

198. Bosch J, Pizcueta P, Feu F, et al. Pathophysiology of portal hypertension. Gastroenterol Clin North Am. 1992;21(1):1–14. PubMed PMID: 1568769. Epub 1992/03/01.

199. Abdelaziz O, Emad-Eldin S, Hussein A, Osman AMA. Role of Doppler ultrasonography in defining normal and abnormal graft hemodynamics after living-donor liver transplant. Exp Clin Transplant. 2017;15(3):306–13. PubMed PMID: 27819194. Epub 2016/11/08.

200. Al-Hamoudi WK. Hemodynamics in the immediate post-transplantation period in alcoholic and viral cirrhosis. World J Gastroenterol. 2010;16(5):608–12.

201. Spitzer AL, Dick AA, Bakhvatsalam R, et al. Intraoperative portal vein blood flow predicts allograft and patient survival following liver transplantation. HPB (Oxford). 2010;12(3):166–73. PubMed PMID: 20590883. Pubmed Central PMCID: PMC2889268. Epub 2010/07/02.

202. Gupta A, Klintmalm GB, Kim PT. Ligating coronary vein varices: an effective treatment of “coronary vein steal” to increase portal flow in liver transplantation. Liver Transpl. 2016;22(7):1037–9. PubMed PMID: 27028766. Epub 2016/03/31.

203. Dalal A. Anesthesia for liver transplantation. Transplant Rev (Orlando). 2016;30(1):51–60. PubMed PMID: 26118926. Epub 2015/06/30.

204. Molmenti EP, Levy MF, Molmenti H, et al. Correlation between intraoperative blood flows and hepatic artery strictures in liver transplantation. Liver Transpl. 2002;8(2):160–3. PubMed PMID: 11862593. Epub 2002/02/28.

205. Kim PT, Saracino G, Jennings L, et al. Ratio of hepatic arterial flow to recipient body weight predicts biliary complications after deceased donor liver transplantation. HPB (Oxford). 2014;16(12):1083–7. PubMed PMID: 25041738. Pubmed Central PMCID: PMC4253331. Epub 2014/07/22.

206. Bonnel AR, Bunchorntavakul C, Rajender Reddy K. Transjugular intrahepatic portosystemic shunts in liver transplant recipients. Liver Transpl. 2014;20(2):130–9. PubMed PMID: 24142390. Epub 2013/10/22.

207. Fagiuoli S, Bruno R, Debernardi Venon W, et al. Consensus conference on TIPS management: techniques, indications, contraindications. Dig Liver Dis. 2017;49(2):121–37. PubMed PMID: 27884494. Epub 2016/11/26.
208. Tripathi D, Therapondos G, Redhead DN, et al. Transjugular intrahepatic portosystemic stent-shunt and its effects on orthotopic liver transplantation. Eur J Gastroenterol Hepatol. 2002;14(8):827–32. PubMed PMID: 12172401. Epub 2002/08/13.

209. Awad N, Horrow MM, Parsikia A, et al. Perioperative management of spontaneous splenorenal shunts in orthotopic liver transplant patients. Exp Clin Transplant. 2012;10(5):475–81. PubMed PMID: 22891944. Epub 2012/08/16.

210. Dua A, McElroy L, Wochinski A, et al. Portal steal syndrome after full-size deceased donor liver transplantation. WMJ. 2016;115(3):147–50. PubMed PMID: 27443092. Epub 2016/07/23.

211. Lee SG, Moon DB, Ahn CS, et al. Ligation of left renal vein for large spontaneous splenorenal shunt to prevent portal flow steal in adult living donor liver transplantation. Transpl Int. 2007;20(1):45–50. PubMed PMID: 17181652. Epub 2006/12/22.

212. Kim B, Kim KW, Song GW, Lee SG. Portal flow steal after liver transplantation. Clin Mol Hepatol. 2015;21(3):314–7. PubMed PMID: 26523275. PubMed Central PMCID: PMC4612294. Epub 2015/11/03.

213. Wu TJ, Dahiya D, Lee CS, et al. Impact of portal venous hemodynamics on indices of liver function and graft regeneration after right lobe living donor liver transplantation. Liver Transpl. 2011;17(9):1035–45. PubMed PMID: 21542130. Epub 2011/05/05.

214. Castillo-Suescun F, Oniscu GC, Hidalgo E. Hemodynamic consequences of spontaneous splenorenal shunts in deceased donor liver transplantation. Liver Transpl. 2011;17(8):891–5. PubMed PMID: 21425432. Epub 2011/03/23.

215. Tang R, Han D, Li M, et al. Left renal vein ligation for large splenorenal shunt during liver transplantation. ANZ J Surg. 2017;87(10):767–72. PubMed PMID: 28851020. Epub 2017/08/30.

216. Kim H, Yoon KC, Lee KW, et al. Tips and pitfalls in direct ligation of large spontaneous splenorenal shunt during liver transplantation. Liver Transpl. 2017;23(7):899–906. PubMed PMID: 28481004. Epub 2017/05/10.

217. Trotter JF. Challenges in living donor liver transplantation. Clin Liver Dis. 2014;18(3):651–60. PubMed PMID: 25017081. Epub 2014/07/16.

218. Graham JA, Samstein B, Emond JC. Early graft dysfunction in living donor liver transplantation and the small for size syndrome. Curr Transplant Rep. 2014;1(1):43–52. PubMed PMID: 27280080. PubMed Central PMCID: PMC4893794. Epub 2014/03/01.

219. Miller CM, Quintini C, Dhawan A, et al. The international liver transplantation society living donor liver transplant recipient guideline. Transplantation. 2017;101(5):938–44. PubMed PMID: 28437386. PubMed Central PMCID: PMC5642345. Epub 2017/04/25.

220. Muzzaile AD, Dagher NN, Montgomery RA, et al. Estimates of early death, acute liver failure, and long-term mortality among live liver donors. Gastroenterology. 2012;142(2):273–80. PubMed PMID: 22108193. Epub 2011/11/24.

221. Abecassis MM, Fisher RA, Oltzoff KM, et al. Complications of living donor hepatic lobectomy – a comprehensive report. Am J Transplant. 2012;12(5):1208–17. PubMed PMID: 22335782. PubMed Central PMCID: PMC3732171. Epub 2012/02/18.

222. Lo CM. Complications and long-term outcome of living liver donors: a survey of 1,508 cases in five Asian centers. Transplantation. 2003;75(3 Suppl):S12–5. PubMed PMID: 12589131. Epub 2003/02/18.

223. Hashikura Y, Ichida T, Umeshita K, et al. Donor complications associated with living donor liver transplantation in Japan. Transplantation. 2009;88(1):110–4. PubMed PMID: 19584689. Epub 2009/07/09.

224. Soejima Y, Shirabe K, Taketomi A, et al. Left lobe living donor liver transplantation in adults. Am J Transplant. 2012;12(7):1877–85. PubMed PMID: 22429497. Epub 2012/03/21.

225. Levy GA, Selzner N, Grant DR. Fostering liver living donor liver transplantation. Curr Opin Organ Transplant. 2016;21(2):224–30. PubMed PMID: 26867047. Epub 2016/02/13.

226. Navasa M, Feu F, Garcia-Pagan JC, et al. Hemodynamic and humoral changes after liver transplantation in patients with cirrhosis. Hepatology. 1993;17(3):355–60. PubMed PMID: 8444409. Epub 1993/03/01.
227. Kelly DM, Demetris AJ, Fung JJ, et al. Porcine partial liver transplantation: a novel model of the "small-for-size" liver graft. Liver Transpl. 2004;10(2):253–63. PubMed PMID: 14762864. Epub 2004/02/06.

228. Demetris AJ, Kelly DM, Eghtesad B, et al. Pathophysiologic observations and histopathologic recognition of the portal hyperperfusion or small-for-size syndrome. Am J Surg Pathol. 2006;30(8):986–93. PubMed PMID: 16861970. Epub 2006/07/25.

229. Tucker ON, Heaton N. The 'small for size' liver syndrome. Curr Opin Crit Care. 2005;11(2):150–5. PubMed PMID: 15758596. Epub 2005/03/11.

230. Herold K, Moser B, Chen Y, et al. Receptor for advanced glycation end products (RAGE) in a dash to the rescue: inflammatory signals gone awry in the primal response to stress. J Leukoc Biol. 2007;82(2):204–12. PubMed PMID: 17513693. Epub 2007/05/22.

231. Soin AS. Smoothing the path: reducing biliary complications, addressing small-for-size syndrome, and making other adaptations to decrease the risk for living donor liver transplant recipients. Liver Transpl. 2012;18(Suppl 2):S20–4. PubMed PMID: 22927168. Epub 2012/08/29.

232. Dahm F, Georgiev P, Clavien PA. Small-for-size syndrome after partial liver transplantation: definition, mechanisms of disease and clinical implications. Am J Transplant. 2005;5(11):2605–10. PubMed PMID: 16212618. Epub 2005/10/11.

233. Selzner M, Kashfi A, Cattral MS, et al. A graft to body weight ratio less than 0.8 does not exclude adult-to-adult right-lobe living donor liver transplantation. Liver Transpl. 2009;15(12):1776–82. PubMed PMID: 19938139. Epub 2009/11/26.

234. Kaido T, Mori A, Ogura Y, et al. Lower limit of the graft-to-recipient weight ratio can be safely reduced to 0.6% in adult-to-adult living donor liver transplantation in combination with portal pressure control. Transplant Proc. 2011;43(6):2391–3. PubMed PMID: 21839274. Epub 2011/08/16.

235. Ben-Haim M, Emre S, Fishbein TM, et al. Critical graft size in adult-to-adult living donor liver transplantation: impact of the recipient's disease. Liver Transpl. 2001;7(11):948–53. PubMed PMID: 11699030. Epub 2001/11/08.

236. Ogura Y, Hori T, El Moghazy WM, et al. Portal pressure <15 mm Hg is a key for successful adult living donor liver transplantation utilizing smaller grafts than before. Liver Transpl. 2010;16(6):718–28. PubMed PMID: 20517905. Epub 2010/06/03.

237. Sainz-Barriga M, Scudeller L, Costa MG, et al. Lack of a correlation between portal vein flow and pressure: toward a shared interpretation of hemodynamic stress governing inflow modulation in liver transplantation. Liver Transpl. 2011;17(7):836–48. PubMed PMID: 21384528. Epub 2011/03/09.

238. Hessheimer AJ, Fondevila C, Taura P, et al. Decompression of the portal bed and twice-baseline portal inflow are necessary for the functional recovery of a “small-for-size” graft. Ann Surg. 2011;253(6):1201–10. PubMed PMID: 21587116. Epub 2011/05/19.

239. Shimamura T, Taniguchi M, Jin MB, et al. Excessive portal venous inflow as a cause of allograft dysfunction in small-for-size living donor liver transplantation. Transplant Proc. 2001;33(1–2):1331. PubMed PMID: 11267312. Epub 2001/03/27.

240. Troisi R, Cammu G, Militerno G, et al. Modulation of portal graft inflow: a necessity in adult living-donor liver transplantation? Ann Surg. 2003;237(3):429–36. PubMed PMID: 12616129. Pubmed Central PMCID: PMC1514313. Epub 2003/03/05.

241. Troisi RI, Berardi G, Tomassini F, Sainz-Barriga M. Graft inflow modulation in adult-to-adult living donor liver transplantation: a systematic review. Transplant Rev (Orlando). 2017;31(2):127–35. PubMed PMID: 27989547. Epub 2016/12/19.

242. Ito T, Kiuchi T, Yamamoto H, et al. Changes in portal venous pressure in the early phase after living donor liver transplantation: pathogenesis and clinical implications. Transplantation. 2003;75(8):1313–7. PubMed PMID: 12717222. Epub 2003/04/30.

243. Sato Y, Yamamoto S, Oya H, et al. Splenectomy for reduction of excessive portal hypertension after adult living-related donor liver transplantation. Hepato-Gastroenterology. 2002;49(48):1652–5. PubMed PMID: 12397756. Epub 2002/10/26.
244. Umeda Y, Yagi T, Sadamori H, et al. Effects of prophylactic splenic artery modulation on portal overperfusion and liver regeneration in small-for-size graft. Transplantation. 2008;86(5):673–80. PubMed PMID: 18791439. Epub 2008/09/16.

245. Troisi R, Ricciardi S, Smeets P, et al. Effects of hemi-portocaval shunts for inflow modulation on the outcome of small-for-size grafts in living donor liver transplantation. Am J Transplant. 2005;5(6):1397–404. PubMed PMID: 15888047. Epub 2005/05/13.

246. Selzner M, Kashfi A, Cattral MS, et al. Live donor liver transplantation in high MELD score recipients. Ann Surg. 2010;251(1):153–7. PubMed PMID: 19858705. Epub 2009/10/28.

247. Hoehn RS, Wilson GC, Wima K, et al. Comparing living donor and deceased donor liver transplantation: a matched national analysis from 2007 to 2012. Liver Transpl. 2014;20(11):1347–55. PubMed PMID: 25044564. Epub 2014/07/22.

248. Olthoff KM, Merion RM, Ghobrial RM, et al. Outcomes of 385 adult-to-adult living donor liver transplant recipients: a report from the A2ALL Consortium. Ann Surg. 2005;242(3):314–23; discussion 23–5. PubMed PMID: 16135918. Pubmed Central PMCID: PMC1357740. Epub 2005/09/02.

249. Olthoff KM, Smith AR, Abecassis M, et al. Defining long-term outcomes with living donor liver transplantation in North America. Ann Surg. 2015;262(3):465–75; discussion 73–5. PubMed PMID: 26258315. Pubmed Central PMCID: PMC4545521. Epub 2015/08/11.

250. Pruett TL, Tibell A, Alabdulkareem A, et al. The ethics statement of the Vancouver Forum on the live lung, liver, pancreas, and intestine donor. Transplantation. 2006;81(10):1386–7. PubMed PMID: 16732173. Epub 2006/05/30.

251. Li C, Wen TF, Yan LN, et al. Risk factors for in-hospital mortality of patients with high model for end-stage liver disease scores following living donor liver transplantation. Ann Hepatol. 2012;11(4):471–7. PubMed PMID: 22700628. Epub 2012/06/16.

252. Perumpail RB, Yoo ER, Cholankeril G, et al. Underutilization of living donor liver transplantation in the United States: Bias against MELD 20 and higher. J Clin Transl Hepatol. 2016;4(3):169–74. PubMed PMID: 27777866. Pubmed Central PMCID: PMC5075001. Epub 2016/10/26.

253. Feng S. Living donor liver transplantation in high model for end-stage liver disease score patients. Liver Transpl. 2017;23(S1):S9–S21. PubMed PMID: 28719072. Epub 2017/07/19.

254. Kaido T, Tomiyama K, Ogawa K, et al. Section 12. Living donor liver transplantation for patients with high model for end-stage liver disease scores and acute liver failure. Transplantation. 2014;97(Suppl 8):S46–7. PubMed PMID: 24849834. Epub 2014/05/23.

255. Liu CL, Fan ST, Lo CM, et al. Live-donor liver transplantation for acute-on-chronic hepatitis B liver failure. Transplantation. 2003;76(8):1174–9. PubMed PMID: 14578749. Epub 2003/10/28.

256. Chok K, Chan SC, Fung JY, et al. Survival outcomes of right-lobe living donor liver transplantation for patients with high model for end-stage liver disease scores. Hepatobiliary Pancreat Dis Int. 2013;12(3):256–62. PubMed PMID: 23742770. Epub 2013/06/08.