

# Expanding Indications of TIPS in the Management of Portal Hypertension Complications

Sai Swarupa Reddy Vulasala<sup>10</sup> Nirmal Kumar Reddy Onteddu<sup>20</sup> Sanjeeva Prasad Kalva<sup>30</sup> Sara Smolinski-Zhao<sup>30</sup>

<sup>1</sup> Department of Radiology, University of Florida College of Medicine, Jacksonville, Florida, United States

<sup>2</sup> Division of Hospital Medicine, Department of Internal Medicine, Flowers Hospital, Dothan, Alabama, United States

<sup>3</sup> Division of Interventional Radiology, Department of Radiology, Massachusetts General Hospital, Boston, Massachusetts, United States

| Clin Interv Radiol ISVIR 2023;7:34–45.

## Abstract

Keywords

- portal hypertension
- ► TIPS
- expanding indications of TIPS

Transjugular intrahepatic portosystemic shunt (TIPS) is a nonsurgical intervention to reduce portal pressure by creating a low-resistance channel between the portal and systemic circulations. It is a well-accepted treatment for gastroesophageal varices and refractory ascites. This review aims to discuss the evidence-based applications of TIPS in other complications of portal hypertension beyond gastroesophageal varices and refractory ascites.

Florida 32209. United States

(e-mail: vulasalaswarupa@gmail.com).

#### Introduction

Transjugular intrahepatic portosystemic shunt (TIPS) is widely used in treating the complications of portal hypertension (PH). PH is defined as an increase in the portal pressure that may lead to the formation of portosystemic collaterals to divert the portal blood to the systemic circulation.<sup>1</sup> PH can be due to structural liver disorders or prehepatic or posthepatic vascular occlusion.<sup>1</sup> Hepatic venous pressure gradient (HVPG) of  $\geq$ 6 mm Hg is diagnostic of PH. Individuals with PH can be asymptomatic or present with ascites, pleural effusion (hepatic hydrothorax [HH]), gastrointestinal (GI) bleeding from variceal hemorrhage, or portal hypertensive gastropathy (PHG), renal failure from hepatorenal syndrome (HRS), and dyspnea from hepatopulmonary syndrome (HPS). These complications develop when the HVPG is  $\geq$  10 mm Hg, termed clinically significant PH.<sup>2</sup> TIPS involves the creation of a conduit between the portal and systemic circulations through the liver parenchyma, thereby reducing the HVPG. Although the technique was introduced in the 1960s, it required additional upgrades to improve patency and mortality rates.<sup>3</sup> To date, many studies described the effectiveness of TIPS in the management of refractory ascites and esophageal varices (EV).<sup>4</sup> In this article, we discussed the evidence in support of other indications of TIPS (**-Table 1**).

Address for correspondence Sai Swarupa Reddy Vulasala, MBBS,

Department of Radiology, University of Florida College of Medicine,

655 West 8th Street, C90, 2nd Floor, Clinical Center, Jacksonville,

#### Hepatic Hydrothorax

HH is defined as the accumulation of transudative fluid (> 500 mL) in the pleural cavity secondary to abdominal ascites. It is prevalent among 5 to 10% of patients with end-stage liver disease.<sup>5,6</sup> The movement of ascitic fluid

article published online May 31, 2022 DOI https://doi.org/ 10.1055/s-0042-1748818. ISSN 1538-8506. © 2022. Indian Society of Vascular and Interventional Radiology. All rights reserved.

This is an open access article published by Thieme under the terms of the Creative Commons Attribution-NonDerivative-NonCommercial-License, permitting copying and reproduction so long as the original work is given appropriate credit. Contents may not be used for commercial purposes, or adapted, remixed, transformed or built upon. (https://creativecommons.org/licenses/by-nc-nd/4.0/)

Thieme Medical and Scientific Publishers Pvt. Ltd., A-12, 2nd Floor, Sector 2, Noida-201301 UP, India

Table 1 Level of evidence of the indications

Indication	Level of evidence
Hepatic hydrothorax	4
Hepatic veno-occlusive disease and Budd–Chiari syndrome	4
Hepatorenal syndrome	4
Portal hypertensive gastropathy	2B
Ectopic varices	4

through the diaphragmatic defects and negative intrathoracic pressure contribute to HH development.<sup>7</sup> HH is seen in the right hemithorax in 85% of cases and in the left among 13% of cases.<sup>8</sup> The presenting symptoms include chest pain and dyspnea on exertion, which worsens with increased fluid accumulation.<sup>8</sup> HH can cause spontaneous bacterial empyema without underlying pneumonia and is observed in 13 to 16% of patients with HH.<sup>9</sup> Diagnosis of HH is by thoracentesis, which demonstrates transudative pleural fluid characteristics as described in **-Table 2**.<sup>7</sup> Management consists of a low sodium diet, diuretics, and therapeutic thoracentesis. Patients requiring repeated thoracentesis every 2 to 3 weeks, albeit on diuretics and sodium-restricted diet, are considered refractory to medical therapy, and they constitute 25% of cases.<sup>5</sup> As such, the definitive therapy is to identify and treat the etiology of ascites through liver transplantation. TIPS can be considered in patients with contraindications to liver transplantation or as a bridge to liver transplantation in patients with refractory HH.<sup>5</sup>

**- Table 3** summarizes the studies involving patients who underwent TIPS for refractory HH. Ditah et al and Campos et al reported that TIPS provided symptomatic relief in three-fourths of the study participants with refractory HH.<sup>5,8</sup> According to Ditah et al and Jindal et al's study, the 45-day mortality rate (17.74%), the 6-

**Table 2** Diagnostic criteria for noninfective hepatichydrothorax

Criteria	Value
Pleural fluid WBC	$< 250/mm^{3}$
Pleural fluid protein	< 2.5 g/dL
Ratio of pleural fluid and serum total protein levels	< 0.5
Ratio of pleural fluid and serum LDH levels	> 0.6
Ratio of pleural fluid and serum albumin levels	> 1.1
Ratio of pleural fluid serum bilirubin levels	< 0.6
рН	> 7.4

Abbreviations: LDH, lactate dehydrogenase; WBC, white blood cell.

month mortality rate (35.9%), and the 1-year survival rate were comparable to the formal indications of TIPS, refractory ascites, and bleeding varices.<sup>5,10</sup> Elderly age group, severe liver disease (Child-Pugh class C, Model for End-Stage Liver Disease [MELD] >15, Child-Turcotte-Pugh [CTP] score >10), elevated creatinine, and lack of response to TIPS are recognized as the predictors of patient mortality.<sup>5</sup> Jindal et al proposed that MELD score >25, spontaneous bacterial peritonitis (SBP), and septic shock are independent predictors of mortality.<sup>10</sup> Although post-TIPS hepatic encephalopathy (HE) (11-66%) was reported in the literature, it was usually responsive to medical therapy without contributing to high mortality rates.<sup>5,8,10</sup> Considering the efficacy of TIPS in HH, its early inclusion may be beneficial as a bridge to definitive treatment.6

#### Budd–Chiari Syndrome

Budd–Chiari syndrome (BCS) is secondary to thrombotic occlusion ranging from the level of the hepatic vein to the

Study	Indication	Sample size	Patient characteristics	Results
Ditah et al <sup>5</sup>	Refractory HH	198	Child class C - 56.9%; Child class B - 40.7%; mean pre-TIPS HVPG - 20.14 mm Hg	CR: 55.8%; PR: 17.6%; AR: 21.2%; 45-day mortality: 17.74%; Overall mortality: 50.17%; HE: 11.7%
Campos et al <sup>8</sup>	Refractory HH	19	Cirrhosis and MELD - 16: 84.2%; Child class C: 47.4%; Child class B: 42.1%	CR: 40%; PR: 33.3%; 30-day mortality: 25%; 1-year mortality: 42.8%; HE: 66.6%
Jindal et al <sup>10</sup>	Refractory HH	51	CTP score: 9.9 ± 1.6; MELD: 18.7 ± 5.4	CR: 20%; PR: 49%; pressure gradient pre- and post-TIPS: $23.1 \pm 3.8$ mm Hg and $7.2 \pm 2.5$ mm Hg; HE: 15%; 6-month mortality rate: 35.9%

**Table 3** Summary of the studies of TIPS in refractory hepatic hydrothorax

Abbreviations: AR, absent response; CR, complete response; CTP score, Child–Turcotte–Pugh score; HE, hepatic encephalopathy; HH, hepatic hydrothorax; HVPG, hepatic venous pressure gradient; MELD, Model for End-Stage Liver Disease; PR, partial response; TIPS, transjugular intrahepatic portosystemic shunt.

right atrium.<sup>11</sup> The incidence of BCS is one in every 2.5 million person-years.<sup>12</sup> Various etiologies of BCS include primary myeloproliferative disorders, hypercoagulable states, oral contraceptive usage and pregnancy, Behcet syndrome, and external compression due to abscess or neoplasms.<sup>13</sup> The classical triad of BCS includes ascites, abdominal pain, and hepatomegaly.<sup>14</sup> The standard management of BCS comprises of anticoagulation and treatment of underlying etiology. An exclusive medical treatment imposed a high mortality rate of 86% in these patients, according to the study by McCarthy et al.<sup>15</sup> An improvement in the survival rate (18% vs. 32%) was reported if the management included thrombolysis, angioplasty, or stent placement.<sup>11</sup> However, Mancuso reported that anticoagulants are the preferred treatment in treating individuals without any signs of PH.<sup>16</sup>

Nonetheless, TIPS is the most common treatment employed in BCS that is complicated by PH. Early TIPS could help prevent the disease progression and hepatic fibrosis, alongside improving the survival outcomes. Liver transplantation is the rescue therapy in individuals with hepatic failure.<sup>17</sup> Critically ill BCS patients awaiting liver transplants may not survive until the surgery and require an emergent procedure such as TIPS to reduce the severity of symptoms. Success rates of TIPS are around 98 to 100% in patients with BCS.<sup>18</sup> The most prevalent indication for TIPS is ascites and variceal bleeding, reported in 100 and 30.9% of cases, respectively.<sup>18</sup> Preprocedural HE and jaundice without hepatic insufficiency are not risk factors for postoperative HE and jaundice. And hence are not considered a contraindication to TIPS.<sup>19,20</sup> Postprocedural complications such as bleeding, HE, and stent malposition were reported to be around 21.4, 2 to 3, and 6%, respectively.<sup>18,21</sup> Seijo et al validated that the treatment of BCS with TIPS can result in a good outcome regardless of the timing of the procedure.<sup>22</sup> In their study of 157 patients, the overall survival rate at 1-, 3-, and 5-year intervals was 88, 83, and 72%, respectively, compared with orthotopic liver transplantation-free survival rates, 85, 78, and 72%, respectively. Based on current evidence, TIPS is highly recommended in early BCS patients as a sole therapy. It might be technically challenging in cases with extensive occlusion of hepatic and suprahepatic veins.<sup>18–22</sup> **- Table 4** summarizes the studies involving patients who underwent TIPS for BCS.

#### **Portal Vein Thrombosis**

Portal vein thrombosis (PVT) is prevalent in 0.7 to 1 individuals for every 100,000 general population.<sup>23</sup> Etiologies for PVT are frequently multifactorial and secondary to myeloproliferative disorders, hepatobiliary malignancies, progressive liver disease, infection, and inflammation.<sup>24</sup> PVT is a common complication of cirrhosis encountered in 20% of patients awaiting liver transplantation.<sup>25</sup> It may manifest with acute or chronic symptoms ranging from asymptomatic to abdominal pain, ascites, variceal bleeding, hypotension, and death.<sup>23</sup> The treatment objective is to resolve symptoms and prevent thrombus extension and secondary complications.<sup>23</sup> As the primary mode of management, anticoagulation has achieved complete recanalization in 53% and partial recanalization in 71% of patients.<sup>26</sup> However, the recanalization rate is lower in patients with thrombus extending to the superior mesenteric vein, chronic PVT, and those with cavernous transformation of portal vein.<sup>25</sup> In addition, around 36% recurrence rate has been reported after the discontinuation of anticoagulation.<sup>27</sup> These limitations of anticoagulation are the determinants that favor TIPS in clinical practice.

Previously, TIPS was contraindicated in patients with PVT due to difficulty identifying the vessels other than collaterals.<sup>28,29</sup> With the advent of contemporary imaging techniques to visualize portal veins, PVT is no longer contemplated as an absolute contraindication to TIPS.<sup>6</sup> In recent times, studies (**-Table 5**) described the effectiveness of TIPS in reducing clot burden, achieving recanalization, and relieving flow stasis. In a recent study by Zhan et al, TIPS improved thrombus burden in 72% of patients while only 27 and 10% of anticoagulated and untreated patients, respectively, demonstrated improvement.<sup>27</sup> In a recent meta-analysis by Valentin et al, the authors reported 84.4% complete and partial recanalization and 74% complete recanalization rates with TIPS.<sup>30</sup> Sun et al demonstrated the efficacy of TIPS in controlling the portal vein pressure and rebleeding rates in patients with chronic PVT (>Table 5). Based on current evidence, TIPS is a feasible treatment to reduce the clot burden and the risk of future portal cavernoma.<sup>30,31</sup> It can be utilized in progressive thrombosis despite anticoagulation and in patients presenting PVT complications such as variceal bleeding.<sup>6</sup>

#### **Hepatorenal Syndrome**

HRS is characterized by renal failure as a result of cirrhosis and PH that meet the International Club of Ascites-Acute Kidney Injury criteria (**► Table 6**).<sup>32,33</sup> It usually develops in patients with decompensated cirrhosis. Previously, HRS was classified into type 1 and type 2 based on serum creatinine. HRS type 1 is defined as a rapidly progressive renal failure in the setting of a precipitating event such as SBP. HRS type 2 comprises slowly progressive renal dysfunction and refractory ascites. According to the International Club of Ascites, HRS type 1 is characterized by serum creatinine > 2.5 mg/dLin < 2 weeks and glomerular filtration rate < 20 mL/min. HRS type 2 is diagnosed when initial serum creatinine is < 2.5mg/dL.<sup>34</sup> Current criteria for HRS include an increase in serum creatinine by > 0.3 mg/dL within 48 hours or by >50% over the baseline within 1 week.<sup>35</sup> An increased hepatic sinusoidal pressure due to cirrhosis leads to systemic vasodilation and vascular underfilling, stimulating renal neurohumoral mechanisms. As a result, sodium and water retention and renal vasoconstriction develop, contributing to HRS.<sup>36</sup> Vasoconstrictor medications along with albumin are the first-line treatment in patients with HRS.<sup>37</sup> Liver transplantation is the standard therapy; however, TIPS can be considered in medically unresponsive patients or in candidates who are unsuitable to transplantation.<sup>37</sup> It

Table 4	Summary of th	e studies of	TIPS in Bu	ıdd–Chiari	syndrome
---------	---------------	--------------	------------	------------	----------

Study	Indication	Sample size	Patient characteristics	Results
Garcia-Pagán et al <sup>11</sup>	BCS	124	Myeloproliferative disorder: 52%; associated IVC thrombus and PV thrombus: 15% and 10%, respectively; mean MELD: 17; refractory ascites: 64%; GI bleed: 15%	1- and 5-year OLT-free survival rate: 88% and 78%, respectively; 5-year OLT-free survival in high-risk patients compared with estimation by Rotterdam BCS index: 71% versus 42%, respectively; TIPS dysfunction: 41%; HE: 21%
Seijo et al <sup>22</sup>	BCS	62	Refractory ascites: 69%; liver failure: 13%; variceal bleeding: 7%	1-, 3-, and 5-year rates of actual survival and OLT-free survival were 88%, 83%, and 72%, and 85%, 78%, and 72%, respectively
Sonavane et al <sup>13</sup>	BCS	42	Mean MELD: 15.38; ascites: 100%; myeloproliferative disease: 40.4%; hyperhomocysteinemia: 12%; all three hepatic vein occlusion; 1005; additional IVC obstruction: 17%	Deaths during follow-up; 26% (36% within 1 month, 18% in 6 months, and 27% in the following period) Causes of death: hematologic disorder: 36%, HE: 27%, intra- abdominal bleed: 18%, and gastrointestinal bleeding: 9% cumulative 1-, 5-, and 10-year OLT-free survival rates were 86%, 81%, and 76%, respectively
Qi et al <sup>65</sup>	BCS	Meta-analysis including 17 studies (each study with > 10 patients)	Refractory ascites; recurrent variceal bleeding	Technical success rates: 91– 100%; pre- and post-TIPS PSG: 27.5 versus 9 mm Hg; clinical improvement: 80– 100%; one- and 5-year cumulative survival rate: 80– 100% and 74–78%, respectively; complications: 0–56%; HE: 0–25%; shunt dysfunction: 18–100% (bare stents: 73%; PTFE stents: 16%)
Fitsiori et al <sup>19</sup>	Refractory BCS	14	BCS-TIPS PI score ≤ 7; chronic myeloproliferative disorder: 57%; hyperhomocysteinemia: 7%; Churg- Strauss syndrome: 7%; paroxysmal nocturnal hemoglobinuria: 7%	Technical success rate: 100%; primary patency: 93%, 85%, 59% at 6, 12, and 24 months, respectively; secondary patency: 100%, 100% and 85% at 6, 12, and 24 months, respectively. TIPS dysfunction: 28.6%
Qi et al <sup>66</sup>	BCS	51 (early TIPS: 19, converted TIPS: 32)	Diffuse hepatic veins obstruction: 23.5%; liver failure: 3%; liver function deterioration: 15.6%; refractory ascites: 19.6%; variceal bleeding: 37.2%	Technical success rate: 100%; portal vein pressure reduced from $28.78 \pm 0.78$ mm Hg to $19.90 \pm 0.77$ mm Hg; HE: 23.5%; shunt dysfunction: 49%; cumulative 1-, 2-, and 3- year rates of being free from shunt dysfunction were 61.6%, $43.9%$ , and $23.4%$ , respectively; 1-, 2-, 3-, 4-, and 5-year survival rates were 83.8%, $81.2%$ , $76.9%$ , $67.3%$ , and $56.09\%$ , respectively

(Continued)

#### Table 4 (Continued)

Study	Indication	Sample size	Patient characteristics	Results
Tripathi et al <sup>20</sup>	BCS	67	MELD: 16.1 $\pm$ 7; CTP score: 8.8 $\pm$ 2.0; hematological risk factors: 78% of patients; ascites: 91%; variceal bleeding; 8.9%	Mean follow-up: 82 months; HE: 15%; primary patency rates were 76% and 27% in covered and uncovered stents; shunt reinterventions were 22% and 100% in covered and uncovered stents; six-, 12-, 24-, 60- and 120-month survival rates were 97%, 92%, 87%, 80%, and 72%, respectively
Fan et al <sup>67</sup>	BCS	60	Ascites: 100%; upper gastrointestinal bleed: 20%; hepatorenal syndrome: 10%; Impaired liver function: 100%; mean CTP score: $9.65 \pm 2.31$ ; proximal ostial occlusion of hepatic vein: 30%; concomitant IVC stenosis: 15%; extensive hepatic vein occlusion; 35%; hepatic venular occlusion: 20%	Technical success: 100%; portal pressure reduced from $41.23 \pm 10.46$ cm H <sub>2</sub> O to $26.68 \pm 6.46$ cm H <sub>2</sub> O; shunt occlusion of intrahepatic portal vein: 5%; hepatic vein reocclusion: 5%
Rosenqvist et al <sup>68</sup>	BCS	13 (from 2003 to 2015)	Hepatomegaly, abdominal pain and ascites: 71%; ascites and fatigue: 21%; unknown clinical presentation: 7%	Technical success rate: 100%; median follow-up period: 3 years; shunt patency: 85% at 1-year and 67% at 2-year follow-up Shunt dysfunction: 30%; HE: 23%; 1- and 5- year OLT-free survival rates were 100% and 93% compared with 47% and 28%, respectively, in 1986–2003

Abbreviations: BCS, Budd–Chiari syndrome; BCS-TIPS PI, BCS-TIPS prognostic index; CTP, Child–Turcotte–Pugh Score; GI, gastrointestinal; HE, hepatic encephalopathy; IVC, inferior vena cava; MELD, Model for End-Stage Liver Disease; OLT, orthotopic liver transplantation; PSG, portosystemic gradient; PTFE, polytetrafluoroethylene; PV, portal vein; TIPS, transjugular intrahepatic portosystemic shunt.

reduces the portal pressure, thereby improving the intravascular volume and cardiac output.

**Table 7** summarizes the studies involving patients who underwent TIPS for HRS. Song et al<sup>37</sup> reported that the 1-year survival rate of HRS-2 and refractory ascitic patients treated with TIPS was 64 and 65%, respectively. Compared with medical management, TIPS improved renal function (52% vs. 83-93%) within a week, and a significant improvement was noticed after 4 weeks. The pooled rate of HE was 49% and was effectively managed with medications. Song et al concluded that TIPS could benefit patients with HRS by improving renal function and survival rates. Charilaou et al<sup>38</sup> conducted a cohort study to compare the efficacy of TIPS and dialysis in HRS patients. They found that the mortality rate was higher in the dialysis-only group compared with the TIPS group (48% vs. 18%). Patients in the TIPS group were three times less likely to be admitted as inpatients than the dialysis-only group (adjusted odds ratio: 0.31; p < 0.001). Shunting with TIPS may impede the progression of renal dysfunction and the need for transplantation. It is more useful in HRS type 2 and could be used to bridge to liver transplantation in medically responsive HRS type 1 individuals. Further studies are required to elaborate on the longterm role of TIPS in HRS patients.

## **Portal Hypertensive Gastropathy**

PHG is seen among 20 to 80% of individuals with PH and described as vascular ectasia of mucosal/submucosal capillaries without any signs of inflammation.<sup>39</sup> The patients with PHG present with acute or chronic GI bleeding that mimics gastric antral vascular ectasia (GAVE). PHG and GAVE have their specific characteristic features on endoscopy, which assists in differentiation. On endoscopy, GAVE displays red spots that can blur together, giving the appearance of a watermelon stomach. PHG appears as a classic mosaic snakeskin-like pattern in the gastric body or fundus progressing to brown or red bulging spots with severity.<sup>39</sup> The severity of PHG correlates with the high CTP score, presence of EV, thrombocytopenia, or splenomegaly. Vasoconstrictors along with resuscitative measures form the mainstay of treatment in acute bleeding. Beta-blockers such as propranolol were evaluated to prevent recurrent bleeding effectively.<sup>39</sup> TIPS is indicated in individuals with recurrent GI

Study	Indication	Sample size	Results
Chen et al <sup>69</sup>	Chronic and completely occluded PVT	18 patients	Mean reduction in PSG from $24.1 \pm 2.3$ mm Hg to $12.1 \pm 3.5$ mm Hg; no complications during the procedure; three deaths during the follow-up period of 16 months due to HCC, severe heart disease, and shunt dysfunction, respectively
Luo et al <sup>28</sup>	Chronic PVT status post-splenectomy	24 patients	Mean reduction in PSG from 22 $\pm$ 4.9 mm Hg to 10.6 $\pm$ 1.6 mm Hg; four HE and five shunt dysfunctions were encountered during a 29-month follow-up
Zhan et al <sup>27</sup>	Nontumoral PVT	52 patients	Thrombus burden improved in 72% of patients treated with TIPS, 27% treated with anticoagulation, and 10% untreated. Complete recanalization was observed in 45% of TIPS patients and in no anticoagulated patients during early follow-up
Valentin et al <sup>30</sup>	PVT	Meta-analysis including 18 studies	Technical success rate: 87%; portal vein recanalization: 84.4%; complete recanalization: 74%; mean change in PSG: 14.5 mm Hg; HE: 25.3%
Sun et al <sup>70</sup>	Chronic PVT and variceal bleeding	189 patients	Technical success rate: 86.2%; mean reduction in portal vein pressure from $27.15 \pm 6.59$ to $19.74 \pm 6.73$ mm Hg; rebleeding rate in TIPS success and fail groups: 15% versus 31%; HE in TIPS success and fail groups: 31% versus 27%; $p = 0.912$

Table 5 Summary of the studies of TIPS in portal vein thrombosis

Abbreviations: HCC, hepatocellular carcinoma; HE, hepatic encephalopathy; PSG, portosystemic gradient; PVT, portal vein thrombosis; TIPS, transjugular intrahepatic portosystemic shunt.

Table 6 ICA-HRS AKI criteria

Diagnosed with cirrhosis or ascites
Diagnosed with AKI based on ICA AKI criteria
Unresponsive to albumin infusion or diuretics withdrawal within 48 hours
No signs of shock
No history of nephrotoxic drug usage
No signs of structural kidney damage such as proteinuria (> 500 mg/dL), microhematuria (> 50 RBCs/HPF), or abnormal ultrasonography

Abbreviations: HPF, high power field; ICA-AKI, International Club of Ascites-Acute Kidney Injury; RBC, red blood cell.

bleeding refractory to  $\beta$ -blockers and iron therapy.<sup>3</sup> The published data are described in **– Table 8**. TIPS has shown significant improvement in the frequency of GI bleeding and transfusion requirements. However, TIPS is found to be

ineffective in controlling symptoms of GAVE. Hence, proper diagnostic differentiation and patient selection are essential among PHG and GAVE.<sup>40,41</sup>

## **Gastric Varices**

Gastric varices (GV) are prevalent in 5 to 33% of cirrhotic and PH patients.<sup>6</sup> They develop at advanced stages of liver disease and constitute 10 to 30% of bleeding episodes with a mortality rate of 45 to 55%.<sup>6,42–44</sup> Studies<sup>45</sup> noted that GV bleeding occurs at lower portosystemic gradient (PSG) compared with EV (17–20 mm Hg vs. 20– 23 mm Hg)<sup>43,45</sup> and hence warrants prompt management. The reason for lower PSG among GV can be explained by its drainage into large caliber gastrorenal shunts in a "downhill" (less resistant caudal flow) process, whereas EVs drain into small-caliber azygos veins in an "uphill" (more resistant cranial flow) process and hence elicit higher PSG.<sup>6,45</sup>

**Table 7** Summary of the studies of TIPS in hepatorenal syndrome

Study	Indication	Sample size	Patient characteristics	Results
Song et al <sup>37</sup>	HRS	128	HRS-1: 77; HRS-2: 51	Short-term, 1-year survival rate in HRS-1 and HRS-2: 72%, 47% and 86%, 64%, respectively; HE: 49%; 1-year mortality in HRS-1 and HRS-2: 0–80% and 31–44%, respectively
Brensing et al <sup>71</sup>	HRS	31	HRS-1: 14; HRS-2: 17	Pre- and post-TIPS PPG: 21 and 13 mm Hg; 3-month survival rate in TIPS and non-TIPS group: 63% and 10%, respectively

Abbreviations: HE, hepatic encephalopathy; HRS, hepatorenal syndrome; PPG, portosystemic pressure gradient; TIPS, transjugular intrahepatic portosystemic shunt.

Study	Indication	Sample size	Patient characteristics	Results
Mezawa et al <sup>40</sup>	PHG	16	Mild PHG: 12; severe PHG: 4	Improved in 4 of 4 severe PHG and 5 of 12 mild PHG patients; pre- and post-TIPS PSG: 23.4 and 14 mm Hg
Kamath et al <sup>41</sup>	РНС	54	Mild PHG: 30; severe PHG: 10; GVE: 14	Endoscopic resolution in 75% of severe PHG and 89% of mild PHG and 0% of GVE patients; 1-year mortality: 50% (27/54); HE: 66%

**Table 8** Summary of the studies of TIPS in portal hypertensive gastropathy

Abbreviations: GVE, gastric vascular ectasia; HE, hepatic encephalopathy; PHG, portal hypertensive gastropathy; PSG, portosystemic gradient; TIPS, transjugular intrahepatic portosystemic shunt.

**Table 9** Theories explaining the suboptimal efficacy of TIPS in gastric varices

Proximity theory	Ineffective decompression due to farther location of GV from the TIPS
Throughput theory	TIPS dysfunction due to supremacy of prominent GRS
Recruitment theory	Emerging of feeding vessels during postembolization of GV period

Abbreviations: GRS, gastrorenal shunt; GV, gastric varices; TIPS, transjugular intrahepatic portosystemic shunt.

Endovascular therapies such as TIPS and balloon-occluded retrograde transvenous obliteration (BRTO) are considered in patients not responding to medical and endoscopic management. TIPS and BRTO can achieve hemostasis in 90 and 95%, respectively, of patients with GV bleed. However, studies reported a lower rebleeding rate (0–20% vs. 25–30%) in the BRTO group compared with the TIPS group.<sup>6</sup> The suboptimal effectiveness of TIPS can be explained by three theories, "proximity," "throughput," and "recruitment" (**- Table 9**).<sup>6</sup>

The incidence of spontaneous portosystemic shunts (SPSs) such as gastrorenal and splenorenal shunts is around 28% in patients with PH.<sup>42</sup> The TIPS placement augments the shunt volume in individuals with SPS and further decreases the portal blood flow (PBF). Reduced PBF enhances the incidence of postinterventional HE, reported by Choi et al, to be around 18%.<sup>42</sup> In contrast, BRTO increases PH by increasing the PBF. Hence, it reduces HE incidence but worsens EV bleeds, ascites, hydrothorax, and PHG.<sup>6,46</sup> Wang et al reviewed the literature comparing TIPS and BRTO procedures. They reported a significant difference in overall survival (risk ratio [RR]: 0.81; p = 0.03) and rebleeding rates (RR: 2.61; p = 0.03) between TIPS and BRTO groups.<sup>47</sup>

In conclusion, the benefits and complications of TIPS and BRTO complement each other, and recently there has been increased application of combined TIPS-BRTO in the management of GV. Implementing BRTO first provides an advantage of increasing the portal vein diameter, making access to TIPS less challenging.<sup>6</sup> The patency of TIPS is also improved with the combined TIPS-BRTO due to the obliteration of competitive SPS.<sup>43</sup>

## **Ectopic Varices**

Ectopic varices can be observed at various abdominal locations such as small bowel, stomas, falciform ligament, biliary tract, vagina, bladder, rectum, umbilicus, and peritoneum.<sup>48</sup> They bleed when the expanding force in varix overcomes the maximum vessel wall tension.<sup>48</sup> Ectopic varices account for 5% of cases presenting with variceal bleed.<sup>49</sup> Management comprises resuscitative measures, vasoconstrictors, endoscopic sclerotherapy, variceal band ligation, transcatheter embolization, or TIPS.<sup>50</sup>

**Table 10** summarizes the studies involving patients who underwent TIPS for ectopic variceal bleed. A study by Oey et al<sup>51</sup> confirmed the efficacy of TIPS in 77% of the patients with ectopic varices, particularly the varices located near enterostomas and associated with mild-moderate liver disease. The rebleeding rate at 1 year was significantly reduced from 39 to 23% due to the usage of expanded polytetrafluoroethylene-covered stents. However, the rebleeding rate of the ectopic varices rate is higher when compared with rebleeding in gastroesophageal varices (94-100% vs. 77%).<sup>51</sup> Increased rebleeding in ectopic varices is attributed to the higher rebleeding rates (50%) in ectopic duodenal varices. The shunt dysfunction is another factor that contributed to rebleeding in three-fourths of the patients and is seen more often in TIPS with a bare-metal stent. Post-TIPS HE was noticed in 30% of patients but manageable with medical treatment or diameter adjustment.

In a study by Kochar et al, TIPS achieved hemostasis in 67% of patients with ectopic varices<sup>48</sup> and 21% presented with rebleeding. According to Vangeli et al, the rebleeding rate was higher in patients who underwent TIPS alone compared with those who underwent a combination of TIPS and variceal embolization (VE) (48% vs. 28%). The patients with rebleeding responded to consecutive VE, and hence Vangeli et al endorsed the inclusion of VE alongside TIPS in the management of ectopic varices.<sup>48,52</sup> However, the routine recommendation of VE needs further studies to demonstrate its efficacy and complications such as propagative thrombus and paradoxical embolization into the systemic circulation.<sup>6</sup>

Study	Indication	Sample size	Patient characteristics	Results
Oey et al <sup>51</sup>	Ectopic variceal bleed	53	Stomal varices: 40%; duodenum: 23%; rectum: 17%; other sites: 20%	Effective in preventing rebleeding: 77% of patients; rebleeding rate at 1, 3, and 5 years: 23%, 26%, and 32%, respectively; HE: 30%
Kochar et al <sup>48</sup>	Ectopic variceal bleed	28	Rectal: 48%; stomal: 28%; duodenal: 14%; other sites: 14%	Portal pressure reduced from $18.2 \pm 6.4$ to $7.2 \pm 3.5$ mm Hg; shunt patency rate at 1, 6, and 9 months: 95%, 89%, and 81%, respectively; survival rate at 1, 3, and 6 months: 81%, 72%, and 61%, respectively; rebleeding: 17%; HE: 30%
Vidal et al <sup>72</sup>	Ectopic variceal bleed	24	Stoma: 33%; duodenal: 20%; ileocolic: 25%; anorectal: 12.5%	Pre- and post-TIPS PSG gradient: 19.7 $\pm$ 5.4 versus 6.4 $\pm$ 3.1 mm Hg, respectively; bleeding resolution: 100%; 1- and 2-year rebleeding and survival rates: 23% and 31%, and 80% and 76%, respectively

Abbreviations: HE, hepatic encephalopathy; PSG, portosystemic gradient; TIPS, transjugular intrahepatic portosystemic shunt.

#### As a Bridge to Liver Transplantation

Liver transplantation is the definitive therapy in liver failure.<sup>53</sup> TIPS can be employed to manage ascites, variceal bleed, and HH in patients awaiting a liver transplant.

Around 14% of patients requiring liver transplantation undergo TIPS placement as a transitory procedure for prompt regulation of illness.<sup>53</sup> **- Table 11** summarizes the studies involving patients who underwent TIPS prior to liver transplantation. Studies of Sellers et al, Valdivieso et al, and Mumtaz et al reported increased intraoperative time, blood transfusion requirement, and length of hospital stay in TIPS patients compared with no-TIPS patients.<sup>53–55</sup> The increased length of hospital stay was due to advanced HE, elderly age, increased cold ischemia time, and MELD scores in a study by Mumtaz et al.<sup>55</sup> However, TIPS allowed patient stabilization and prolonged the waiting time between TIPS intervention and transplant surgery.<sup>53</sup> Graft survival rates, mortality rate, and retransplant rates in TIPS patients were noted to be similar to those in no-TIPS patients.53,55

## **TIPS after Liver Transplantation**

Recurrence of liver disease can lead to the development of PH in patients with liver transplants.<sup>56</sup> TIPS procedure is challenging in liver transplant recipients due to changes in the liver anatomy. Lerut et al pointed the difficulty of cannulating the graft hepatic and portal veins during the TIPS procedure in individuals who underwent piggyback cavo-caval anastomoses.<sup>57</sup>

• **Table 12** summarizes the studies involving patients who underwent TIPS after liver transplantation. Chen et al<sup>56</sup> studied patients who underwent liver transplantation and were experiencing refractory ascites, variceal hemorrhage, and HH. They reported a 98% technical success rate with a resolution of symptoms in 57% of refractory ascites, 69% of variceal bleeding, and 56% of HH patients. However, 33% of patients experienced HE, 16% required shunt revision, and 19% required retransplantation. Chen et al concluded that it was reasonable to suggest TIPS in liver transplant recipients if they develop recurrent PH. In contrast, the technical success rate was 68.2% in a study by King et al. They also

Study	Indication	Sample size	Patient characteristics	Results
Mumtaz et al <sup>55</sup>	TIPS prior to liver transplantation	1366		TIPS increased the waiting time for transplant (408 $\pm$ 553 days) compared with no-TIPS (183 $\pm$ 330 days); no significant effect of TIPS was noted on mortality and retransplant rate
Amesur et al <sup>73</sup>	TIPS prior to liver transplantation	12	Variceal bleeding: 50%; ascites: 50%	Child A patients had superior survival; two patients with ascites experienced death within 1 week due to liver failure

Table 11 Summary of the studies of TIPS prior to liver transplantation

Abbreviation: TIPS, transjugular intrahepatic portosystemic shunt.

Study	Indication	Sample size	Patient characteristics	Results
Chen et al <sup>56</sup>	TIPS after liver transplantation	213	Refractory ascites: 78%; variceal hemorrhage: 17%; hydrothorax: 4%	Technical success: 98%; success rates of TIPS after OLT in patients with refractory ascites, variceal hemorrhage, and hydrothorax were 57%, 69%, and 56%; HE: 33%; 30-day mortality rate and 1-year survival rate: 11% and 53%, respectively; subsequent retransplantation: 19%
King et al <sup>58</sup>	TIPS after liver transplantation	22 transplanted patients (cases) and 44 nontransplants (controls)	Variceal bleeding: 36.4%; refractory ascites: 63.6%	Pre- versus post-TIPS PSG in cases and controls: 21.0 versus 9.9 mm Hg and 22.4 versus 6.9 mm Hg, respectively. technical success rates in cases and controls: 68.2 and 95.5%; clinical success rates: 77.2 versus 93.2%, respectively
Lerut et al <sup>57</sup>	TIPS after liver transplant	8	Refractory ascites: 62; HH and ascites: 12.5%; bleeding esophageal varices: 12.5%; repeated biliary surgery: 12.5%	Technical success rate: 1,005; complete response: 37.5%; partial response: 37.5%; unfavorable: 25%

Table 12 Summary of the studies of TIPS after liver transplantation

Abbreviations: HE, hepatic encephalopathy; HH, hepatic hydrothorax; OLT, orthotopic liver transplantation; PSG, portosystemic gradient; TIPS, transjugular intrahepatic portosystemic shunt.

inferred that TIPS was not beneficial in patients with MELD > 15 (hazard ratio [HR] = 5.846), and retransplant could be considered in such individuals.<sup>58</sup>

#### **TIPS Prior to an Abdominal Surgery**

Extrahepatic abdominal surgeries in patients with chronic liver disease (CLD) are associated with a considerable postoperative mortality rate of 10 to 76%.<sup>59</sup> The CTP score allows the prediction of postoperative mortality, with Child–Pugh C being the worst prognostic factor and Child–Pugh A and B carrying poor outcomes. Ascites in CLD hinders wound healing, resulting in wound dehiscence and infections. Alongside, increased portal pressure leads to complications such as intraoperative bleeding and hepatic decompensation.<sup>59</sup> Preoperative TIPS lessens the portal pressure, thereby reducing the complications.<sup>6</sup>

**- Table 13** summarizes the studies involving patients who underwent TIPS prior to extrahepatic abdominal surgery. Tabchouri et al reported an 85% operability rate in patients undergoing preoperative TIPS placement.<sup>60</sup> In 2006, Vinet et al concluded that preoperative TIPS had not demonstrated beneficial postoperative effects.<sup>61</sup> In support of their study, Tabchouri et al described that although TIPS reduces postoperative ascites (HR = 0.3), it worsens MELD (HR = 2.3), and there is no statistically significant effect on 90-day mortality rates (HR = 0.720; 0.180–2.920) and complications (HR: 0.670; 0.270–1.680).<sup>60</sup> In patients with HVPG > 13 mm Hg, increased intraoperative blood transfusion requirement (HR: 4.1) and increased postoperative sepsis (HR: 2.8) were reported.<sup>60</sup> Based on current evidence,<sup>60,61</sup> TIPS in the

Study	Indication	Sample size	Patient characteristics	Results
Vinet et al <sup>61</sup>	Extrahepatic abdominal surgery	18	Antrectomy: 5; colectomy: 10; small bowel resection: 1; pancreatectomy: 1; nephrectomy: 1	No significant improvement in mortality, complications were reported
Tabchouri et al <sup>60</sup>	Extrahepatic abdominal surgery	With TIPS: 66; without TIPS: 68	Colorectal surgery: 68; upper Gl and pancreatic surgery: 13; hernia and incisional hernia: 17; cholecystectomy: 13; other: 13	Operability rate: 85%; postoperative ascites hazard ratio: 0.330; similar mortality and complications such as bleeding in TIPS and no-TIPS group

**Table 13** Summary of the studies of TIPS in extrahepatic abdominal surgery

Abbreviations: GI, gastrointestinal; TIPS, transjugular intrahepatic portosystemic shunt.

Study	Indication	Sample size	Patient characteristics	Results
Tsauo et al <sup>74</sup>	HPS	12	Moderate HPS: 16%; severe HPS: 16%; very severe HPS: 66%	75% of patients had improved oxygenation; pre- and post-TIPS PSG: 18.2 mm Hg versus 6.5 mm Hg; 22% patients had recurrence despite patent shunt

**Table 14** Summary of the studies of TIPS in hepatopulmonary syndrome

Abbreviations: HPS, hepatopulmonary syndrome; PSG, portosystemic gradient; TIPS, transjugular intrahepatic portosystemic shunt.

preoperative management plan of extrahepatic abdominal surgeries is not recommended. Still, further studies need to be conducted to establish the efficacy of TIPS.

#### Hepatopulmonary Syndrome

HPS constitutes impaired blood oxygenation due to dilated intrapulmonary vasculature secondary to hepatic cirrhosis.<sup>62</sup> It is observed among 5 to 32% of patients with liver disease.<sup>63,64</sup> The patients present with platypnea, pathognomonic of HPS, and dyspnea.<sup>62</sup> Diagnostic criteria include the presence of hepatic disease, alveolar arterial oxygen gradient (A-a O2) of  $\geq$  15 mm Hg ( $\geq$  20 mm Hg in patients above 64 years old), and dilated pulmonary vasculature demonstrated on contrast-enhanced echocardiography (bubble study). Although liver transplantation is the definitive treatment for HPS, the application of TIPS is increasing due to its effective reduction in portal pressure, which in turn lowers the vasodilators causing pulmonary vascular constriction.<sup>63</sup> In addition, the mortality rate is reported to be around 16 to 33% in HPS patients undergoing liver transplantation.<sup>64</sup>

Left portal vein (LPV) TIPS improves symptoms of HPS better than right portal vein TIPS, and the incidence of HE is diminished in LPV TIPS.<sup>6</sup> A study by Tsauo et al (**-Table 14**)<sup>64</sup> reported statistically significant improvement in A-a O2 and oxygenation after 1 month of TIPS creation. However, the recovery is transient, and they observed worsening hypoxemia 3 months after TIPS. Hence, the study concluded that TIPS cannot be performed as a solitary treatment for HPS but could be considered a bridge to definitive treatment.<sup>64</sup>

#### Conclusion

TIPS has been widely applied to manage complications of PH. Besides its routine clinical applications—refractory ascites and EV—TIPS has gained paramount importance in varied conditions, as outlined in this article. Further prospective studies would enhance the strength of evidence and recommendations for these indications.

#### **Conflict of Interest**

S.P.K. reports grants from NIH, BD, Black Swan, and Trisalus for Institution; reports royalties from Elsevier, Springer, and Thieme for himself; reports consulting fees from Penumbra, Okami Medical, Boston Scientific, Medtronic, Covidien, US Vascular, Dova Pharmaceuticals, Instylla, and BD for himself; reports payment from Stony Brook University, American Institute of Biology, UT Houston, and NACCME for himself; reports payment for expert testimony from Southern Institute for Medical and Legal Affairs LLC for himself; reports participation from NIH for institution; reports leadership from Chief, Interventional Radiology, Massachusetts General Hospital, Boston, MA: Chair, Vascular Panel, ACR Appropriateness Criteria; International Editor, Journal of Clinical Interventional Radiology ISVIR; Assistant Editor, Radiology - Cardiothoracic, RSNA; reports stock from Biogen Inc, Clover Health Investments Corp, Inovio Pharmaceuticals, Moderna Inc, Pfizer Inc, Novavax Inc, Orphazyme, Cassava Sciences Inc, Vivos Therapeutics Inc, Ardelyx Inc, Althea Health, Sarepta Therapeutics, Clover Health Investments Corp, CureVac BV, Immunoprecise Antibodies Ltd, Infinity Pharmaceuticals Inc, Zymergen Inc, BioNTech SE, Trillium Therapeutics Inc, Theravance Biopharma Inc, Doximity Inc, Eargo Inc, Allogent Therapeutics Inc, NRx Pharmaceuticals Inc, Atea pharmaceuticals Inc, for himself and spouse; and reports financial or nonfinancial interests as Adjunct Associate Professor from University of Texas Southwestern Medical Center.

#### References

- 1 Pinchot JW, Kalva SP, Majdalany BS, et al; Expert Panels on Interventional Radiology and Vascular Imaging. ACR Appropriateness Criteria® radiologic management of portal hypertension. J Am Coll Radiol 2021;18(55):S153–S173
- 2 Augustin S, Millán L, González A, et al. Detection of early portal hypertension with routine data and liver stiffness in patients with asymptomatic liver disease: a prospective study. J Hepatol 2014; 60(03):561–569
- 3 Tripathi D, Stanley AJ, Hayes PC, et al. Transjugular intrahepatic portosystemic stent-shunt in the management of portal hypertension. Gut 2020;69(07):1173–1192
- 4 Smith M, Durham J. Evolving indications for tips. Tech Vasc Interv Radiol 2016;19(01):36–41
- 5 Ditah IC, Al Bawardy BF, Saberi B, Ditah C, Kamath PS. Transjugular intrahepatic portosystemic stent shunt for medically refractory hepatic hydrothorax: a systematic review and cumulative metaanalysis. World J Hepatol 2015;7(13):1797–1806
- 6 Rajesh S, George T, Philips CA, et al. Transjugular intrahepatic portosystemic shunt in cirrhosis: an exhaustive critical update. World J Gastroenterol 2020;26(37):5561–5596
- 7 Garbuzenko DV, Arefyev NO. Hepatic hydrothorax: an update and review of the literature. World J Hepatol 2017;9(31):1197–1204
- 8 Campos S, Gomes D, Sofia C. Transjugular intrahepatic portosystemic shunt in refractory hydrothorax - a contribution to an unexplored indication. Eur J Gastroenterol Hepatol 2016;28(06): 661–666
- 9 Chow E, Khiatah B, Frugoli A. Refractory spontaneous bacterial empyema in cirrhotic patient. Case Rep Gastrointest Med 2021 2021:6685998

- 10 Jindal A, Mukund A, Kumar G, Sarin SK. Efficacy and safety of transjugular intrahepatic portosystemic shunt in difficult-tomanage hydrothorax in cirrhosis. Liver Int 2019;39(11): 2164–2173
- Garcia-Pagán JC, Heydtmann M, Raffa S, et al; Budd-Chiari Syndrome-Transjugular Intrahepatic Portosystemic Shunt Group. TIPS for Budd-Chiari syndrome: long-term results and prognostics factors in 124 patients. Gastroenterology 2008;135(03): 808–815
- 12 Martens P, Nevens F. Budd-Chiari syndrome. United European Gastroenterol J 2015;3(06):489–500
- 13 Sonavane AD, Amarapurkar DN, Rathod KR, Punamiya SJ. Long term survival of patients undergoing TIPS in Budd-Chiari syndrome. J Clin Exp Hepatol 2019;9(01):56–61
- 14 Ryu RK, Durham JD, Krysl J, et al. Role of TIPS as a bridge to hepatic transplantation in Budd-Chiari syndrome. J Vasc Interv Radiol 1999;10(06):799–805
- 15 McCarthy PM, van Heerden JA, Adson MA, Schafer LW, Wiesner RH. The Budd-Chiari syndrome. Medical and surgical management of 30 patients. Arch Surg 1985;120(06):657–662
- 16 Mancuso A. An update on the management of Budd-Chiari syndrome: the issues of timing and choice of treatment. Eur J Gastroenterol Hepatol 2015;27(03):200–203
- 17 Lupasco I, Dumbrava V-T. Diagnosis and therapy of Budd Chiari syndrome. Med Pharm Rep 2021;94(Suppl No 1):S68–S71
- 18 Inchingolo R, Posa A, Mariappan M, et al. Transjugular intrahepatic portosystemic shunt for Budd-Chiari syndrome: a comprehensive review. World J Gastroenterol 2020;26(34): 5060–5073
- 19 Fitsiori K, Tsitskari M, Kelekis A, Filippiadis D, Triantafyllou K, Brountzos E. Transjugular intrahepatic portosystemic shunt for the treatment of Budd-Chiari syndrome patients: results from a single center. Cardiovasc Intervent Radiol 2014;37(03):691–697
- 20 Tripathi D, Macnicholas R, Kothari C, et al. Good clinical outcomes following transjugular intrahepatic portosystemic stent-shunts in Budd-Chiari syndrome. Aliment Pharmacol Ther 2014;39(08): 864–872
- 21 Hayek G, Ronot M, Plessier A, et al. Long-term outcome and analysis of dysfunction of transjugular intrahepatic portosystemic shunt placement in chronic primary Budd-Chiari syndrome. Radiology 2017;283(01):280–292
- 22 Seijo S, Plessier A, Hoekstra J, et al; European Network for Vascular Disorders of the Liver. Good long-term outcome of Budd-Chiari syndrome with a step-wise management. Hepatology 2013;57 (05):1962–1968
- 23 Sharma AM, Zhu D, Henry Z. Portal vein thrombosis: when to treat and how? Vasc Med 2016;21(01):61–69
- 24 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(01):42–51
- 25 Rodrigues SG, Sixt S, Abraldes JG, et al. Systematic review with meta-analysis: portal vein recanalisation and transjugular intrahepatic portosystemic shunt for portal vein thrombosis. Aliment Pharmacol Ther 2019;49(01):20–30
- 26 Loffredo L, Pastori D, Farcomeni A, Violi F. Effects of anticoagulants in patients with cirrhosis and portal vein thrombosis: a systematic review and meta-analysis. Gastroenterology 2017;153(02): 480–487.e1
- 27 Zhan C, Prabhu V, Kang SK, et al. Comparison of non-tumoral portal vein thrombosis management in cirrhotic patients: TIPS versus anticoagulation versus no treatment. J Clin Med 2021;10 (11):2316
- 28 Luo J, Li M, Zhang Y, et al. Percutaneous transhepatic intrahepatic portosystemic shunt for variceal bleeding with chronic portal vein occlusion after splenectomy. Eur Radiol 2018;28(09): 3661–3668

- 29 Ferrusquía-Acosta J, Hernández-Gea V. TIPS indications and contraindications—pushing the limits: is earlier better? Curr Hepatol Rep 2019;18(01):87–95
- 30 Valentin N, Korrapati P, Constantino J, Young A, Weisberg I. The role of transjugular intrahepatic portosystemic shunt in the management of portal vein thrombosis: a systematic review and meta-analysis. Eur J Gastroenterol Hepatol 2018;30(10): 1187–1193
- 31 Keshava SN, Moses V, Sharma A, et al. Technical and mediumterm clinical outcomes of transjugular intrahepatic portosystemic shunt with fluoroscopy and additional trans-abdominal ultrasound guidance. Indian J Radiol Imaging 2021;31(04): 858–866
- 32 Guevara M, Ginès P, Bandi JC, et al. Transjugular intrahepatic portosystemic shunt in hepatorenal syndrome: effects on renal function and vasoactive systems. Hepatology 1998;28(02): 416-422
- 33 Amin AA, Alabsawy EI, Jalan R, Davenport A. Epidemiology, pathophysiology, and management of hepatorenal syndrome. Semin Nephrol 2019;39(01):17–30
- 34 Arroyo V, Ginès P, Gerbes AL, et al. Definition and diagnostic criteria of refractory ascites and hepatorenal syndrome in cirrhosis. International Ascites Club. Hepatology 1996;23(01):164–176
- 35 Angeli P, Ginès P, Wong F, et al; International Club of Ascites. Diagnosis and management of acute kidney injury in patients with cirrhosis: revised consensus recommendations of the International Club of Ascites. Gut 2015;64(04):531–537
- 36 Rössle M, Gerbes AL. TIPS for the treatment of refractory ascites, hepatorenal syndrome and hepatic hydrothorax: a critical update. Gut 2010;59(07):988–1000
- 37 Song T, Rössle M, He F, Liu F, Guo X, Qi X. Transjugular intrahepatic portosystemic shunt for hepatorenal syndrome: a systematic review and meta-analysis. Dig Liver Dis 2018;50(04):323–330
- 38 Charilaou P, Devani K, Petrosyan R, Reddy C, Pyrsopoulos N. Inpatient mortality benefit with transjugular intrahepatic portosystemic shunt for hospitalized hepatorenal syndrome patients. Dig Dis Sci 2020;65(11):3378–3388
- 39 Ripoll C, Garcia-Tsao G. The management of portal hypertensive gastropathy and gastric antral vascular ectasia. Dig Liver Dis 2011;43(05):345–351
- 40 Mezawa S, Homma H, Ohta H, et al. Effect of transjugular intrahepatic portosystemic shunt formation on portal hypertensive gastropathy and gastric circulation. Am J Gastroenterol 2001;96 (04):1155–1159
- 41 Kamath PS, Lacerda M, Ahlquist DA, McKusick MA, Andrews JC, Nagorney DA. Gastric mucosal responses to intrahepatic portosystemic shunting in patients with cirrhosis. Gastroenterology 2000;118(05):905–911
- 42 Choi YH, Yoon CJ, Park JH, Chung JW, Kwon JW, Choi GM. Balloonoccluded retrograde transvenous obliteration for gastric variceal bleeding: its feasibility compared with transjugular intrahepatic portosystemic shunt. Korean J Radiol 2003;4(02):109–116
- 43 Lipnik AJ, Pandhi MB, Khabbaz RC, Gaba RC. Endovascular treatment for variceal hemorrhage: TIPS, BRTO, and combined approaches. Semin Intervent Radiol 2018;35(03):169–184
- 44 Sarin SK, Lahoti D, Saxena SP, Murthy NS, Makwana UK. Prevalence, classification and natural history of gastric varices: a longterm follow-up study in 568 portal hypertension patients. Hepatology 1992;16(06):1343–1349
- 45 Morrison JD, Mendoza-Elias N, Lipnik AJ, et al. Gastric varices bleed at lower portosystemic pressure gradients than esophageal varices. J Vasc Interv Radiol 2018;29(05):636–641
- 46 Kim SK, Lee KA, Sauk S, Korenblat K. Comparison of transjugular intrahepatic portosystemic shunt with covered stent and balloon-occluded retrograde transvenous obliteration in managing isolated gastric varices. Korean J Radiol 2017;18 (02):345–354

- 47 Wang ZW, Liu JC, Zhao F, et al. Comparison of the effects of TIPS versus BRTO on bleeding gastric varices: a meta-analysis. Can J Gastroenterol Hepatol 202020205143013
- 48 Kochar N, Tripathi D, McAvoy NC, Ireland H, Redhead DN, Hayes PC. Bleeding ectopic varices in cirrhosis: the role of transjugular intrahepatic portosystemic stent shunts. Aliment Pharmacol Ther 2008;28(03):294–303
- 49 Norton ID, Andrews JC, Kamath PS. Management of ectopic varices. Hepatology 1998;28(04):1154–1158
- 50 Akhter NM, Haskal ZJ. Diagnosis and management of ectopic varices. Gastrointestinal Intervention 2012;1(01):3–10
- 51 Oey RC, de Wit K, Moelker A, et al. Variable efficacy of TIPSS in the management of ectopic variceal bleeding: a multicentre retro-spective study. Aliment Pharmacol Ther 2018;48(09):975–983
- 52 Vangeli M, Patch D, Terreni N, et al. Bleeding ectopic varicestreatment with transjugular intrahepatic porto-systemic shunt (TIPS) and embolisation. J Hepatol 2004;41(04):560–566
- 53 Sellers CM, Nezami N, Schilsky ML, Kim HS. Transjugular intrahepatic portosystemic shunt as a bridge to liver transplant: current state and future directions. Transplant Rev (Orlando) 2019;33(02):64–71
- 54 Valdivieso A, Ventoso A, Gastaca M, et al. Does the transjugular intrahepatic portosystemic influence the outcome of liver transplantation? Transplant Proc. 2012;44(06):1505–1507
- 55 Mumtaz K, Metwally S, Modi RM, et al. Impact of transjugular intrahepatic porto-systemic shunt on post liver transplantation outcomes: study based on the United Network for Organ Sharing database. World J Hepatol 2017;9(02):99–105
- 56 Chen B, Wang W, Tam MD, Quintini C, Fung JJ, Li X. Transjugular intrahepatic portosystemic shunt in liver transplant recipients: indications, feasibility, and outcomes. Hepatol Int 2015;9(03): 391–398
- 57 Lerut JP, Goffette P, Molle G, et al. Transjugular intrahepatic portosystemic shunt after adult liver transplantation: experience in eight patients. Transplantation 1999;68(03):379–384
- 58 King A, Masterton G, Gunson B, et al. A case-controlled study of the safety and efficacy of transjugular intrahepatic portosystemic shunts after liver transplantation. Liver Transpl 2011;17(07): 771–778
- 59 Boike JR, Flamm SL. Transjugular intrahepatic portosystemic shunts: advances and new uses in patients with chronic liver disease. Clin Liver Dis 2020;24(03):373–388
- 60 Tabchouri N, Barbier L, Menahem B, et al. Original study: transjugular intrahepatic portosystemic shunt as a bridge to abdominal surgery in cirrhotic patients. J Gastrointest Surg 2019;23(12): 2383–2390
- 61 Vinet E, Perreault P, Bouchard L, et al. Transjugular intrahepatic portosystemic shunt before abdominal surgery in cirrhotic patients: a retrospective, comparative study. Can J Gastroenterol 2006;20(06):401–404

- 62 Grilo-Bensusan I, Pascasio-Acevedo JM. Hepatopulmonary syndrome: what we know and what we would like to know. World J Gastroenterol 2016;22(25):5728–5741
- 63 Wallace MC, James AL, Marshall M, Kontorinis N. Resolution of severe hepato-pulmonary syndrome following transjugular portosystemic shunt procedure. BMJ Case Rep 2012;2012: bcr0220125811
- 64 Tsauo J, Zhao H, Zhang X, et al. Effect of transjugular intrahepatic portosystemic shunt creation on pulmonary gas exchange in patients with hepatopulmonary syndrome: a prospective study. J Vasc Interv Radiol 2019;30(02):170–177
- 65 Qi X, Yang M, Fan D, Han G. Transjugular intrahepatic portosystemic shunt in the treatment of Budd-Chiari syndrome: a critical review of literatures. Scand J Gastroenterol 2013;48(07):771–784
- 66 Qi X, Guo W, He C, et al. Transjugular intrahepatic portosystemic shunt for Budd-Chiari syndrome: techniques, indications and results on 51 Chinese patients from a single centre. Liver Int 2014;34(08):1164–1175
- 67 Fan X, Liu K, Che Y, et al. Good clinical outcomes in Budd–Chiari syndrome with hepatic vein occlusion. Dig Dis Sci 2016;61(10): 3054–3060
- 68 Rosenqvist K, Sheikhi R, Eriksson L-G, et al. Endovascular treatment of symptomatic Budd-Chiari syndrome - in favour of early transjugular intrahepatic portosystemic shunt. Eur J Gastroenterol Hepatol 2016;28(06):656–660
- 69 Chen Y, Ye P, Li Y, Ma S, Zhao J, Zeng Q. Percutaneous transhepatic balloon-assisted transjugular intrahepatic portosystemic shunt for chronic, totally occluded, portal vein thrombosis with symptomatic portal hypertension: procedure technique, safety, and clinical applications. Eur Radiol 2015;25(12):3431–3437
- 70 Sun X-Y, Wang G-C, Wang J, Huang G-J, Zhang C-Q. Transjugular intrahepatic portosystemic shunt is effective in patients with chronic portal vein thrombosis and variceal bleeding. Hepatobiliary Pancreat Dis Int 2021;20(02):128–136
- 71 Brensing KA, Textor J, Perz J, et al. Long term outcome after transjugular intrahepatic portosystemic stent-shunt in nontransplant cirrhotics with hepatorenal syndrome: a phase II study. Gut 2000;47(02):288–295
- 72 Vidal V, Joly L, Perreault P, Bouchard L, Lafortune M, Pomier-Layrargues G. Usefulness of transjugular intrahepatic portosystemic shunt in the management of bleeding ectopic varices in cirrhotic patients. Cardiovasc Intervent Radiol 2006;29(02):216–219
- 73 Amesur NB, Zajko AB, Orons PD, Sammon JK, Casavilla FA. Transjugular intrahepatic portosystemic shunt in patients who have undergone liver transplantation. J Vasc Interv Radiol 1999;10 (05):569–573
- 74 Tsauo J, Weng N, Ma H, Jiang M, Zhao H, Li X. Role of transjugular intrahepatic portosystemic shunts in the management of hepatopulmonary syndrome: a systemic literature review. J Vasc Interv Radiol 2015;26(09):1266–1271