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**Case Control Study**

Covered transjugular intrahepatic portosystemic stent-shunt vs large volume paracentesis in patients with cirrhosis: A real-world propensity score-matched study

Dhaliwal A et al. Covered TIPSS vs LVP

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**Abstract**

**BACKGROUND**

Refractory ascites has a 1-year survival rate of 50%. In selected patients, treatment options include liver transplantation (LT) or transjugular intrahepatic portosystemic stent shunt (TIPSS).

**AIM**

To assess the outcomes of patients who underwent a TIPSS compared to large volume paracentesis (LVP).

**METHODS**

Retrospective study of patients who underwent a covered TIPSS or LVP for refractory or recurrent ascites over 7 years. Primary outcome was transplant-free survival (TFS). Further analysis was done with propensity score matching (PSM).

**RESULTS**
There were 150 patients [TIPSS group (n = 75), LVP group (n = 75)]. Seven patients in
the TIPSS group underwent LT vs 22 patients in the LVP group. Overall median follow
up, 20 (0.47-179.53) mo. In the whole cohort, there was no difference in TFS [hazard
ratio (HR): 0.80, 95% confidence interval (CI): 0.54-1.21]; but lower de novo hepatic
encephalopathy with LVP (HR: 95%CI: 0.20-0.96). These findings were confirmed
following PSM analysis. On multivariate analysis albumin and hepatocellular
carcinoma at baseline were associated with TFS.

CONCLUSION
Covered TIPSS results in similar TFS compared to LVP in cirrhotic patients with
advanced liver failure. Liver transplant assessment should be considered in all potential
candidates for TIPSS. Further controlled studies are recommended to select appropriate
patients for TIPSS.

Key Words: Portal hypertension; Liver cirrhosis; Transjugular intrahepatic
portosystemic shunt; Ascites; Large volume paracentesis

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Core Tip: This retrospective, propensity score matched study comparing covered TIPSS
vs LVP in patients with cirrhosis has shown that transplant free survival was similar.

INTRODUCTION
Liver cirrhosis is a significant cause of morbidity and mortality in the United Kingdom
and worldwide[1-3]. Clinically significant portal hypertension leads to decompensation,
with ascites often the first evidence of hepatic decompensation[4]. Ascites is initially
treated with diuretics however for those with diuretic intractable ascites, frequent large-
volume paracentesis with albumin cover is the remaining option. Patients with refractory ascites have a 1-year survival rate of 50%. In a selection of these patients, treatment options include liver transplantation and transjugular intrahepatic portosystemic stent shunt (TIPSS)\(^5\).

Ascites occurs due to two main mechanisms: portal hypertension and sodium and water retention. Liver cirrhosis alters the normal hepatic architecture by progressive collagen deposition (fibrosis) and nodular regeneration within the hepatocytes and distortion of the hepatic vasculature\(^6\). As a result, all these changes, hepatic sinusoids have a reduction in their compliance and there is an increase in resistance to portal flow. Splanchnic vasodilation, mediated by nitrous oxide (extra-hepatic production increases in cirrhosis), and soluble guanylyl cyclase dependent protein kinase G signalling, and other vasoactive mediators, contributes to hyperdynamic circulation manifested as increased cardiac output and heart rate, with a decreased systemic vascular resistance and a low arterial blood pressure. This leads to greater blood flow through the portal vein, which in presence of increased resistance, contributes to portal hypertension. Clinically significant portal hypertension, (defined as a hepatic venous portal gradient > 10 mmHg)\(^7\), whereby there is an increase in the hydrostatic pressure within hepatic sinusoids, compounds this. Hence there is further transudation of fluid into the peritoneal cavity and subsequent ascites\(^8\). The hyperdynamic circulation results in reduced central blood volume due to splanchnic vasodilation. This triggers increases in renal sympathetic activity including activation of renin-angiotensin and aldosterone systems, and antidiuretic hormone to improve central blood volume. This enhances sodium reabsorption within the renal tubules and collecting ducts resulting in increased sodium and water retention\(^9\)-\(^11\).

The only curative treatment option in refractory ascites is liver transplantation. This is suitable for selected patients through a rigorous screening and assessment process\(^12\)-\(^14\). Whilst diuretic therapy and large volume paracenteses provide a therapeutic approach, more definitive treatment with TIPSS is an option in selected patients\(^14\)-\(^16\). TIPSS reduces portal pressure with an initial increase in cardiac output, right atrial
pressure, and pulmonary artery pressure leading to a secondary reduction in systemic vascular resistance and effective arterial blood volume. These haemodynamic changes have been reported to return to pre TIPSS level with time. Additionally, there is an increase in urinary sodium excretion and glomerular filtration rate\[^{13,17-20}\]. It is well established that covered TIPSS is superior to non-covered TIPSS with significantly reduced stent dysfunction and recurrence of portal hypertension-related complications\[^{21,22}\]. Patient selection for TIPSS has several considerations including the severity of liver disease, renal function, vascular anatomy, nutritional status, risk factors for hepatic encephalopathy (HE), and cardiac function\[^{23}\]. Patient selection is paramount to a beneficial outcome. One of the most important and disabling complications of TIPSS is the development of de novo HE which occurs in 30%-50% of patients\[^{13}\].

For those with diuretic intractable ascites, and who are unsuitable for TIPSS and liver transplantation, frequent palliative large volume paracentesis (LVP) remains the main course of symptom management. These patients often have end-stage liver disease, with high short-term mortality, so LVP remains a safe and effective management strategy\[^{5,8}\]. There is recent interest in long term abdominal drains, which may reduce the need for hospital visits but can be complicated by increased infections. Further research is required\[^{14}\].

This study aims to assess the outcome of those who underwent covered TIPSS vs LVP for the treatment of refractory ascites. We aimed to ascertain the following: (1) transplant-free survival (TFS); (2) Clinical or biochemical variables that predict survival; and (3) Risk factors for developing HE.

**MATERIALS AND METHODS**

**Study design**

A retrospective cohort study was performed at the Liver Unit, Queen Elizabeth Hospital, University Hospitals Birmingham. Our study groups consisted of two groups. Group 1 comprised patients who underwent a covered TIPSS between April 2010 to
November 2017. Group 2 (the standard of care group) comprised patients who underwent frequent LVP with albumin cover between January 2011 to November 2017. Patients were identified using our institute’s electronic information technology informatics system and electronic patient database through coding methods.

We included patients with liver cirrhosis, and an age greater than or equal to 18. For Group 1, we included those who had covered TIPSS for refractory or recurrent ascites as an elective procedure. For Group 2 we included those who had > 1 LVP per month.

We excluded patients who did not have a diagnosis of liver cirrhosis, those who underwent a TIPSS insertion for variceal haemorrhage, those who underwent frequent paracentesis due to malignancy including hepatocellular carcinoma (HCC) with malignant spread or underwent liver transplantation before first LVP.

The primary outcome measure of this study was TFS. We had several secondary endpoints which included effectiveness of TIPSS compared with LVP as quantified by the number of LVP per month and complications of TIPSS in the form of de novo HE. Follow up was carried out until 2017 or until death had occurred.

**Treatments**

**TIPSS group:** After the patient’s informed consent, TIPSS (Viatorr® stent-graft; GORE, Flagstaff, AR, United States) was placed in the standard method described previously, and the tract between right hepatic and portal veins was dilated to 8-10 mm\(^{23}\). Pre-TIPSS (either portal pressure or hepatic venous pressure gradient) and post-TIPSS pressures were measured. Patients were monitored for any immediate complications for at least 48 h post-procedure. None of these patients received anticoagulation post-TIPSS. The patency was routinely assessed by follow up 6-12 mo doppler US at clinic review and venography if indicated by the US scan or clinical deterioration.

**LVP group:** LVP was performed according to the local guidelines with 20 g of albumin administered for every 2 L to 3 L ascites fluid drained, in concordance with the local
and national guidance. The procedures were performed as a day case for most patients. All patients were followed up routinely in the outpatient clinic at 3-6 mo intervals.

Parameters

We compared biochemical and clinical parameters for each group. The clinical parameters were obtained from clinical medical electronic records. These included mortality rates, complications of portal hypertension (HE and gastric or oesophageal varices), development of HCC, and use of non-selective beta-blockers (NSBB). We collected laboratory records to assess severity staging in the form of model for end-stage liver disease (MELD) and United Kingdom MELD, liver function, hematology and renal function parameters. These were recorded at the time of the index intervention [i.e. TIPSS or first LVP (1 ± 1 d)].

Statistics

Categorical variables such as gender were expressed as a number and percentage. Numerical data were expressed as a mean ± standard deviation for normally distributed data. Data was also expressed as median with a range where appropriate.

Comparisons between the two groups were performed using an unpaired t test or chi-squared test. Both univariate and multivariable analyses were used to control for differences in selected independent parameters such as MELD score. A Cox regression analysis was used to identify clinical and biochemical variables predicting survival. Actuarial probability survival curves were constructed using the Kaplan-Meier method and compared with the log-rank test. To confirm the validity of the results in matched cohorts, a propensity score analysis was performed. Propensity score matching (1:1) with matched tolerance of 0.02 was performed to account for covariates (platelet count, MELD, gender, sodium and age). Further supplemental sensitivity analysis was done using the propensity score weighting method[24]. Statistical significance was established at a P < 0.05. SPSS statistical software (version 27) was used to perform the analyses.
RESULTS

Patient characteristics

There were 106 patients identified as receiving a covered TIPSS. We excluded 31 patients due to other indications such as a repeated procedure with an existing TIPSS in situ, and who did not fulfil the inclusion criteria. Thus, the TIPSS group comprised 75 patients. There were 89 patients with liver cirrhosis who underwent frequent LVP with albumin cover within our hospital, however, 14 were excluded as they met the exclusion criteria. Hence, the LVP group comprised of n = 75 (Figure 1, Table 1).

With regards to the severity of liver disease, compared to the TIPSS group, the LVP group had a significantly higher mean United Kingdom MELD (51.5 ± 4.2 vs 54.6 ± 4.8) and MELD (11.5 ± 3.9 vs 15.9 ± 5.3) (Table 2). There was no difference in the use of NSBB, presence of varices, HCC, or history of spontaneous bacterial peritonitis (SBP) at baseline.

Table 1 also shows the characteristics of the propensity score-matched cohort. The TIPSS group and LVP group comprised 40 patients each. The baseline characteristics were well matched although fewer patients underwent liver transplantation and had refractory ascites in the TIPSS group. These differences were also present in the PSM cohort but to a lesser degree. Table 2 shows that laboratory data and clinical scoring systems were similar in both groups.

Clinical outcomes during follow up

Effectiveness of TIPSS: The portal pressure gradient (PPG) decreased from 15.7 ± 4.9 mmHg to 6.7 ± 2.7 mmHg following TIPSS implantation with a mean PPG reduction of 54.7% ± 17.6 %. In 59 and 16 patients the stent was dilated to 8 mm and 10 mm respectively. TIPSS resulted in a significant reduction in the requirement for paracentesis per month compared with the LVP group (0.1 ± 0.6 vs 1.2 ± 0.6, \( P < 0.001 \)). Diuretics were required during the clinical course in all patients in the LVP group, and in only 14.7% of patients in the TIPSS group. Further LVP was not required in 74.7% of patients in the TIPSS group. There was no difference in the baseline aetiology of liver
disease ($P = 0.44$), MELD ($P = 0.69$), CPS ($P = 0.24$), bilirubin ($P = 0.05$), platelets ($P = 0.53$), albumin ($P = 0.98$), age ($P = 0.96$), gender ($P = 0.39$), history of SBP ($P = 0.6$), presence of varices ($P = 0.24$), TIPSS diameter ($P = 0.30$) and PPG % reduction post TIPSS ($P = 0.80$), PPG post TIPSS ($P = 0.58$) between those who did or did not require further LVP post TIPSS.

**Transplantation free survival**

**Whole cohort:** The actuarial rate of TFS at 6 mo, 12 mo, 24 mo and 60 mo for each group is as follows: TIPSS group 76%, 64%, 53%, 20%; LVP group 78%, 55%, 36%, 15%, respectively. Figure 2 shows the Kaplan Meier curve to represent this. The causes of death are detailed in Table 3. Analysis using log-rank statistics did not reveal any significant difference in TFS (HR: 1.24, 95%CI: 0.83-1.86). In the TIPSS arm, an increased number of paracentesis per month (HR: 1.35, 95%CI: 1.02-1.77) and 8 mm stent diameter (HR: 2.93, 95%CI: 1.31-6.52) was associated with a worse TFS.

Univariate analysis demonstrated that albumin, HCC at baseline and CPS predicted TFS (Table 4). Multivariable analyses showed that only HCC at baseline and albumin were significant as predictors of survival (Tables 4 and 5). Further analysis excluding patients with HCC at baseline, which can cause significant confounding, demonstrated that there remained no significant differences in TFS.

**Propensity score-matched cohort:** The actuarial rate of TFS at 6 mo, 12 mo, 24 mo and 60 mo for each group is as follows: TIPSS group 66%, 50%, 41%, and 17%; LVP group 81%, 54%, 34% respectively (Figure 3). There was no significant difference in TFS (HR: 1.00, 95%CI: 0.58-1.73). A sensitivity analysis using propensity score weighting method confirmed the lack of significance between the cohorts (HR: 0.95, 95%CI: 0.60-1.53).

**Hepatic encephalopathy**

**Whole cohort:** The actuarial rate of *de novo* HE at 6 mo, 12 mo, and 24 mo for each group are as follows; TIPSS: 28%, 28%, 31%. LVP group 5%, 7%, and 16% (Figure 4).
This was a statistically significant difference in favour of LVP (HR: 0.44, 95%CI: 0.20-0.96). In the TIPSS arm, neither diameter of the stent (P = 0.35), PPG post-TIPSS (P = 0.88), or degree of PPG reduction (0.74) influenced the rate of de novo HE. TIPSS reduction was performed successfully in two patients due to refractory HE.

**Propensity score-matched cohort:** The actuarial rate of de novo HE at 6 mo, 12 mo, and 24 mo for each group are as follows; TIPSS: 33%, 33%, 33%. LVP group 5%, 11%, and 11% (Figure 5). This was a statistically significant difference in favour of LVP [Figure 5; HR: 0.30 95%CI: 0.10-0.94, P = 0.03 (log-rank)]. A sensitivity analysis using a propensity score weighting method confirmed the difference in favour of LVP (HR: 0.38, 95%CI: 0.15-0.95).

**Liver transplantation**

Seven patients in the TIPSS group underwent liver transplantation vs twenty-two patients in the LVP group (P = 0.002). The mean time to transplantation during follow up was 13.4 ± 16.5 mo, with no difference between the groups.

**DISCUSSION**

We have shown in our retrospective study of 150 patients, which is one of the largest comparative series in the literature, that TFS following covered TIPSS for refractory ascites is similar to LVP, with increased HE with TIPSS. We also found that the control of ascites was significantly better with TIPSS.

We controlled for confounders by using a propensity score matching, which corroborated these findings. We found that the long-term outcomes are indeed poor in both groups, with 5-year TFS in TIPSS and LVP cohorts of 20% and 15% respectively. This compares with 5-year survival post-liver transplantation in Europe of 71%.[25]. This supports and reinforces the view that liver transplantation is the best option in eligible patients with end stage liver disease (ESLD).
There is only one randomized controlled trial of covered TIPSS vs LVP which showed improved TFS without increased risk of HE\textsuperscript{[26]}. This trial only included patients with recurrent ascites. We include both recurrent and refractory ascites and did not find the presence of recurrent ascites influenced TFS. The impact on TFS remains uncertain based on a recent network meta-analysis of 287 participants which showed that TIPSS resulted in greater resolution of ascites compared to LVP but no difference in mortality or adverse events\textsuperscript{[27]}. Our real-world cohort reinforces these findings. A recent retrospective study of TIPSS vs LVP showed that TIPSS was not independently associated with TFS.

Hepatic encephalopathy is one of the symptoms that significantly affect those with advanced liver failure. It can manifest both covertly and overtly in patients. Whilst a thorough assessment for the presence of HE is required before consideration for TIPSS, it remains a challenge to manage. We found that \textit{de novo} HE rates were higher with TIPSS in the whole cohort and the cohort after excluding TIPSS contraindications. Bucsics \textit{et al.}\textsuperscript{[28]} also found in their retrospective study of TIPSS vs LVP similar rates of \textit{de novo} HE in both LVP and TIPSS cohorts.

A recent study concluded that covered TIPSS resulted in \textit{superior control of ascites} without increasing the risk for overt HE as compared to LVP\textsuperscript{[28]}. We believe our data reflects the real-world experience, and has the strength of a larger sample size and follow up. There is interest in the role of rifaximin prophylactically before TIPSS in reducing the risk of HE after TIPSS\textsuperscript{[29]}. We did not use rifaximin to prevent HE post TIPSS, as the evidence of benefit was published after the recruitment period for our study. Controlled expansion stents can also reduce the risk of passive dilatation of stents and HE but further controlled data is required\textsuperscript{[30]}.

We only had 7 patients (9\%) of our TIPSS cohort undergo liver transplantation, which is far less than in the LVP cohort where 22 (29\%) patients underwent transplantation. This suggests that patients with TIPSS may have been less likely to be referred for liver transplantation due to control of ascites. Furthermore, the current scoring methods of liver disease are not as helpful in refractory ascites. Our data would
strongly suggest that all patients undergoing TIPSS must be considered for transplant at an early stage, in particular, those not responding to TIPSS or where there is deteriorating liver function.

Serum albumin and hepatocellular carcinoma emerged as independent predictors of survival in keeping with recently published data\cite{28}. However, there were differences in the rate of HCC at baseline which were not controlled by propensity score matching, unlike our study. We also performed a separate analysis excluding HCC and found no differences in TFS.

An important finding of a recent study was the superior control of ascites post TIPSS with early TIPSS insertion at lower paracentesis frequencies and creatinine levels. Persistent ascites post-TIPSS was a predictor of liver transplantation and death\cite{31}. We found that lack of effect of TIPSS, as judged by the increased need for LVP post TIPSS was associated with poor TFS. This would suggest that patients with poor efficacy after TIPSS should be considered for liver transplantation at an early stage.

The diameter of the stent is an important consideration. There is conflicting literature on the impact of stent diameter on outcomes, and no recommendation can be made at this time\cite{14}. In most of our patients, the stents were dilated to 8 mm, and interestingly the use of 10 mm diameter was associated with better TFS. We would advise caution in the interpretation of this finding due to the small numbers of patients with 10 mm stents. We did not find the post TIPSS PPG or proportional reduction of PPG post TIPSS associated with TIPSS efficacy, and this was also the case in the recent study by Piecha et al\cite{31}. It is also worth noting that passive dilatation of TIPSS stent occurs even with 8 mm dilatation, although recent controlled expansion stents are much less prone to this phenomenon\cite{30,32}. Therefore, the stent diameter and PPG at the time of TIPSS insertion is likely to change significantly over time.

The recognition of sarcopenia is an evolving consideration in those with ESLD. Sarcopenia is associated with increased mortality in those with ESLD and nutritional assessment is now recommended in patients considered for elective TIPSS\cite{33}. However, the data for patients with sarcopenia and advanced cirrhosis and undergoing TIPSS is
inconsistent. A recent study found that sarcopenia (defined as muscle mass alone) did not have an impact on survival in a similar cohort of patients with refractory ascites undergoing TIPSS\textsuperscript{34}. It is important to recognise the lack of functional elements in this definition of sarcopenia as this may demonstrate different outcomes. Frailty, which incorporates this functional element\textsuperscript{35} which should also be a consideration in future work. Whilst we did not comment on the degree of sarcopenia in our cohort, it may have been a contributing factor and should be a future consideration to research.

Our study does have some limitations. The retrospective nature introduces selection bias, but we selected consecutive patients in a real-world setting from a single institution, and the large sample size is a major strength. Moreover, we carefully controlled for key confounders using propensity score matching and still retained a total of 80 patients. The increased rate of transplantation in the LVP group also introduces bias concerning competing risks. However, we selected TFS to minimise this bias. PSM resulted in similar follow up for TIPSS and LVP groups which could help to minimise this confounder.

**CONCLUSION**

We found that after controlling for confounding factors, our retrospective real-world data shows that TFS was similar following covered TIPSS for refractory or recurrent ascites compared with LVP. The presence of HCC, low albumin, and poor response to TIPSS are associated with poor survival. The long term outcomes following TIPSS are poor. From this, we can conclude that liver transplantation must be considered for refractory ascites in selected patients. For patients not suitable for liver transplantation, other interventions for refractory ascites could be considered palliative. Further prospective studies are required in multicentre controlled trials to identify prognostic markers to aid patient selection for interventions for refractory ascites.
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