Long term outcome after transjugular intrahepatic portosystemic stent-shunt in non-transplant cirrhotics with hepatorenal syndrome: a phase II study


Abstract

Background—Recent small studies on hepatorenal syndrome (HRS) indicate some clinical benefit after transjugular intrahepatic portosystemic stent-shunt (TIPS) but sufficient long term data are lacking.

Aim—We studied prospectively feasibility, safety, and long term survival after TIPS in 41 non-transplantable cirrhotics with HRS (phase II study).

Patients and methods—HRS was diagnosed using current criteria (severe (type I) HRS, n=21; moderate (type II) HRS, n=20). Thirty one patients (14 type I, 17 type II) received TIPS (8–10 mm) while advanced liver failure excluded shunting in 10. During follow up (median 24 months) we analysed renal function and survival (Kaplan-Meier).

Results—TIPS markedly reduced the portal pressure gradient (21 (5) to 13 (4) mm Hg (mean (SD)); p<0.001) with one procedure related death (3.2%). Renal function deteriorated without TIPS but improved (p<0.001) within two weeks after TIPS (creatinine clearance 18 (15) to 48 (42) ml/min; sodium excretion 9 (16) to 77 (78) mmol/24 hours) and stabilised thereafter. Following TIPS, three, six, 12, and 18 month survival rates were 81%, 71%, 48%, and 35%, respectively. As only 10% of non-shunted patients survived three months, total survival rates were 63%, 56%, 39%, and 29%, respectively. Multivariate Cox regression analysis revealed bilirubin (p<0.001) and HRS type (p<0.05) as independent survival predictors after TIPS.

Conclusions—TIPS provides long term renal function and probably survival benefits in the majority of non-transplantable cirrhotics with HRS. These data warrant controlled trials evaluating TIPS in the management of HRS.

Keywords: hepatorenal syndrome; transjugular intrahepatic portosystemic stent-shunt; liver cirrhosis; portal hypertension; ascites; renal failure

Hepatorenal syndrome (HRS) is a common and severe complication of advanced liver cirrhosis. After the onset of HRS, patients have only a minimal chance of renal functional recovery and a poor prognosis under medical therapy. Nine out of 10 patients with advanced HRS die within 10 weeks, most within the first month after diagnosis. To date, the only established therapy that guarantees long term improvement in renal function and prolongation of survival is timely liver transplantation. However, most of these patients are non-transplant candidates at the time of HRS diagnosis but require intensive medical management which may include haemodialysis in some patients. Furthermore, transplant organ shortage limits urgent transplant even for transplantable patients.

Patients with refractory ascites who are at high risk of HRS can be effectively treated by insertion of a transjugular intrahepatic portosystemic stent-shunt (TIPS). However, data on recovery of renal function after TIPS in these patients are controversial. One study reported an increase in glomerular filtration rates (GFR) in six month survivors. In another small randomised trial comparing TIPS with large volume paracentesis for refractory ascites, GFR improved only marginally after TIPS while natriuresis increased significantly. As refractory ascites and HRS share a similar pathophysiology, TIPS has been tried as a rescue measure in patients with advanced HRS. So far, preliminary short term data are favourable. However, these series are small (1–7 severe HRS patients) and often lack follow up data beyond three months. Furthermore, HRS was often defined differently and included patients who were candidates for transplant rescue. This hampers outcome analysis, especially for those high risk cirrhotic patients who are actually not transplant candidates at the time of HRS diagnosis.

The present phase II study provides prospective long term data on HRS outcome following TIPS as rescue treatment in a large cohort of cirrhotics in whom liver transplantation was contraindicated at the time of HRS diagnosis.

Patients and methods

STUDY POPULATION AND DEFINITION OF HRS
Between January 1995 and November 1998 we evaluated 45 consecutive patients with ad-

Abbreviations used in this paper: TIPS, transjugular intrahepatic portosystemic stent-shunt; HRS, hepatorenal syndrome; GFR, glomerular filtration rates; EF1, endothelin 1; PRA, plasma renin activity; AR, active renin.
Advanced cirrhosis of the liver, refractory ascites, and recently detected renal insufficiency. Forty one of these patients (25 males, 16 females) were excluded from live transplantation as a therapeutic option and were enrolled in the present study of non-surgical management (table 1). The cause of liver cirrhosis was alcoholism in 31 patients and chronic viral hepatitis in eight. Two patients had cryptogenic cirrhosis. Within two months prior to detection of renal insufficiency, 22 patients (54%) had suffered clinical complications related to portal hypertension, such as upper gastrointestinal haemorrhage (12 patients) and spontaneous bacterial peritonitis (10 patients). These complications had been effectively treated and none of these patients had a relapse within at least two weeks prior to diagnosis of renal failure. At presentation, 26 alcoholics were still active drinkers and nine of these had suffered recent severe alcohol abuse and clinically suspected concomitant acute alcoholic hepatitis (liver biopsy was performed in two and confirmed the clinical diagnosis). Thirty one patients were considered eligible for TIPS as an ultimate rescue treatment whereas 10 (25%) were excluded from TIPS because of one or more persistent contraindications (bilirubin levels >15 mg/dl (five), Child-Pugh points >12 (four), or spontaneous severe encephalopathy (three)).

All patients were studied after a minimum of five days without diuretics on a 50 mmol/day sodium diet as part of their oral nutrition. Oliguric patients had a bladder catheter inserted during diagnostic work up. Assessment of renal function was based on at least 2–3 samples of daily urine volume and serum parameters obtained prior to individual treatment assignment. All patients fulfilled recent consensus criteria for HRS,2 which are, in brief: proven liver cirrhosis with severe ascites; no recent nephrotoxic medication or severe fluid losses; no renal abnormalities on ultrasound; no nephritic sediment; proteinuria less than 500 mg/day; no signs of active infection or other reasons for renal impairment; and finally no improvement in renal function after withdrawal of diuretics and appropriate volume expansion (200 ml of 20% albumin and/or 1–2 litres of isotonic saline). HRS was graded as HRS type I (severe and rapidly progressive: serum creatinine >2.5 mg/dl or creatinine clearance <20 ml/min; >50% loss in clearance within two weeks) or HRS type II (moderate and stable or slowly progressive: serum creatinine >1.5 mg/dl or creatinine clearance <40 ml/min). Overall, we diagnosed 21 patients with type I and 20 with type II HRS. Fourteen patients with type I HRS (seven of whom required haemodialysis due to persistent oligoanuria with life threatening pulmonary oedema and hyperkalaemia) and 17 with type II HRS received TIPS treatment. The median interval from the first detection of renal insufficiency to TIPS insertion was shorter in type I HRS (2.2 weeks; range 0.3–6) than in type II HRS (4.0 weeks; range 1.5–5) but mean values were not significantly different (3.0 (1.6) v 4.3 (0.3) weeks). Among the 10 non-stented patients, seven had type I and three had type II

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Table 1 Clinical baseline characteristics of non-transplant cirrhotics with hepatorenal syndrome (HRS) (n=41)

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Total (n=41)</th>
<th>Non-TIPS (n=10)</th>
<th>TIPS (n=31)</th>
<th>p Value</th>
<th>HRS type I (n=21)</th>
<th>HRS type II (n=20)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (y)</td>
<td>57 (11)</td>
<td>50 (15)</td>
<td>59 (9)</td>
<td>&lt;0.05</td>
<td>56 (9)</td>
<td>61 (6)</td>
<td>NS</td>
</tr>
<tr>
<td>Aetiology of LC</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Alcoholic</td>
<td>31</td>
<td>6</td>
<td>25</td>
<td></td>
<td>13</td>
<td>18</td>
<td>0.02</td>
</tr>
<tr>
<td>Post-hepatitic</td>
<td>8</td>
<td>4</td>
<td>4</td>
<td></td>
<td>3</td>
<td>3</td>
<td>NS</td>
</tr>
<tr>
<td>Cryptogenic</td>
<td>2</td>
<td>0</td>
<td>2</td>
<td>NS</td>
<td>0</td>
<td>2</td>
<td>NS</td>
</tr>
<tr>
<td>Causes for transplant refusal</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Active alcoholism</td>
<td>26</td>
<td>6</td>
<td>20</td>
<td></td>
<td>13</td>
<td>13</td>
<td>11</td>
</tr>
<tr>
<td>Advanced age</td>
<td>5</td>
<td>1</td>
<td>4</td>
<td></td>
<td>4</td>
<td>1</td>
<td>NS</td>
</tr>
<tr>
<td>Advanced cachexia</td>
<td>9</td>
<td>3</td>
<td>6</td>
<td></td>
<td>7</td>
<td>2</td>
<td>NS</td>
</tr>
<tr>
<td>Recent cancer</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>NS</td>
<td>1</td>
<td>0</td>
<td>NS</td>
</tr>
<tr>
<td>Child-Pugh score</td>
<td>10.4 (1.9)</td>
<td>12.8 (1.0)</td>
<td>9.5 (1.4)</td>
<td>&lt;0.001</td>
<td>11.0 (2.1)</td>
<td>9.7 (1.5)</td>
<td>0.02</td>
</tr>
<tr>
<td>Child-Pugh grade</td>
<td>A</td>
<td>0</td>
<td>0</td>
<td>NS</td>
<td>0</td>
<td>0</td>
<td>NS</td>
</tr>
<tr>
<td>C</td>
<td>16</td>
<td>6</td>
<td>10</td>
<td>&lt;0.01</td>
<td>16</td>
<td>4</td>
<td>NS</td>
</tr>
<tr>
<td>Bilirubin (mg/dl)</td>
<td>6.8 (9.0)</td>
<td>17.4 (12.0)</td>
<td>3.1 (3.3)</td>
<td>&lt;0.01</td>
<td>10.6 (11.7)</td>
<td>4.9 (12.7)</td>
<td>NS</td>
</tr>
<tr>
<td>Albumin (g/dl)</td>
<td>4.0 (1.9)</td>
<td>4.0 (1.1)</td>
<td>4.0 (1.1)</td>
<td>NS</td>
<td>4.0 (1.1)</td>
<td>4.0 (1.1)</td>
<td>NS</td>
</tr>
<tr>
<td>Serum creatinine (mg/dl)</td>
<td>1.9 (1.9)</td>
<td>1.9 (1.9)</td>
<td>1.9 (1.9)</td>
<td>NS</td>
<td>1.9 (1.9)</td>
<td>1.9 (1.9)</td>
<td>NS</td>
</tr>
<tr>
<td>Serum sodium (mmol/l)</td>
<td>130 (60)</td>
<td>130 (60)</td>
<td>130 (60)</td>
<td>NS</td>
<td>130 (60)</td>
<td>130 (60)</td>
<td>NS</td>
</tr>
<tr>
<td>Oliguria (&lt;25 ml/24 h)</td>
<td>481 (365)</td>
<td>286 (268)</td>
<td>449 (352)</td>
<td>&lt;0.05</td>
<td>385 (434)</td>
<td>568 (279)</td>
<td>&lt;0.05</td>
</tr>
</tbody>
</table>

*p values were calculated from subgroup comparisons of clinical baseline characteristics.

To convert values for creatinine to µmol/l, multiply by 88.4; to convert values for urea to mmol/l, multiply by 0.167; to convert values for bilirubin to µmol/l, multiply by 17.1.
HRS. Preliminary short term data on changes in renal function up to six months after TIPS treatment in 16 of our patients (six with type I and 10 with type II HRS) have been reported previously.13

All patients were considered ineligible for liver transplantation for the following reasons (table 1): active alcoholism (n=26), eligibility for major surgery (age over 65 years or advanced cachexia) (n=14), and recent oropharyngeal cancer (n=1). Six active alcoholics and four patients considered poor candidates for major surgery were excluded from TIPS and received the best medical support, including adequate volume/electrolyte management, paracentesis and albumin substitution, and vasoconstrictive drugs, as indicated.2 Patients gave written informed consent for TIPS as a salvage procedure as well as for portal and systemic blood sampling according to the study protocol which was approved by the local ethics committee.

TIPS INSERTION

The day before the scheduled TIPS insertion all patients received a therapeutic paracentesis followed by intravenous administration of albumin (8 g/l ascites volume). After an overnight fast and bec-tst, patients were taken to the radiology department for the TIPS procedure. We punctured the portal system under ultrasound guidance using a 9 French liver biopsy needle (Cook, Germany) and established small diameter (8–10 mm) stent-shunts either with Palmaz stents or Wallstents, as described in other TIPS reports.15–17 Our target was a reduction in portal pressure gradient (portal minus vena cava superior pressure) of 30–50%. In patients with no or only mild ascites after paracentesis this gradient differs only minimally from the more commonly assessed gradient of portal minus vena cava inferior pressure.18 For technical reasons, to minimise the amount of contrast media as well as the duration of the procedure, we used the modified gradient to assess the relative pressure changes induced by TIPS. The initial technical success rate was 100% without apparent adverse effects during the TIPS procedure. Patients received a single dose of antibiotic prophylaxis of cefuroxime (1.5 g), and mida-zolam (5–15 mg) with pethidine (75–150 mg) for anaesthesia. Intravenous heparin was given for prevention of shunt thrombosis (bolus dose of 2500–5000 U followed by constant infusion for 1–2 weeks, targeted at an activated partial thromboplastin time of 60–80 seconds). TIPS patency was monitored by repeated Doppler ultrasound (twice weekly within the first two weeks) and routine radiography 10–17 days after TIPS in all but two patients. Thereafter, Doppler ultrasound was performed every 2–3 months or whenever clinically indicated. Additional radiography was performed only for suspected TIPS dysfunction. TIPS patency was maintained during the study period in all patients. Following a minimum of one week after TIPS, diuretics were gradually reintroduced (spironolactone 50–200 mg/day and frusemide 20–80 mg/day) if daily spontaneous diuresis and 40–60% reduction of the previous ascites volume had not occurred. Excretion of sodium and potassium was guided using serum electrolyte and creatinine levels. Thromboplastin time of 60–80 seconds. TIPS was inserted in 20 patients (11 type I and nine type II HRS) from the portal vein (prior to stenting) and from the systemic circulation to determine endothelin-1 (ET-1, ELISA assay, reference values 0.73 (0.03) pmol/l) levels and activation of the renin-angiotensin system. The latter was estimated by measuring plasma renin activity (PRA) in 12 (RIA assay, reference values: baseline <5 ng/ml/3 hours, stimulated <10 ng/ml/3 hours) or by active renin (AR; RIA assay, reference values: baseline <25 ng/l, stimulated <50 ng/l) in the last eight patients. A second pair of portal and systemic plasma samples were obtained during the first invasive TIPS evaluation after 12–15 days.

STATISTICAL ANALYSIS

Follow up parameters were compared with baseline values using two sided Student’s t tests or by Mann-Whitney U test, as appropriate, for patients at risk at a given time interval after confirmation of changes over time by MANOVA. Categorical data were analysed with Fisher’s exact test. All values are given as mean (SD). Differences at p<0.05 were considered significant. Estimation of survival was based on the Kaplan-Meier method and on the log rank test for group comparisons. Cox regression models were used for multivariate analysis of outcome predictors. SPSS-PC+ software was used on an IBM computer.

Results

STUDY POPULATION

Baseline clinical characteristics in the total study population (table 1) were similar for type I and type II HRS patients with respect to age and aetiology of liver cirrhosis, but type I HRS patients showed a more advanced degree of liver dysfunction, as assessed by routine parameters. Interestingly, despite significantly different glomerular filtration rate parameters, there was a similar reduction in serum sodium and sodium excretion in type I and type II patients.

RENAL FUNCTION AFTER TIPS INSERTION

Thirty one patients received TIPS (table 1) on average 3.4 (1.3) weeks after detection of renal insufficiency, which reduced the portal pres-
TIPS for hepatorenal syndrome

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§§Portal v

VASOCONSTRICTIVE MEDIATORS

Mean baseline systemic ET-1 was elevated 2–3-fold above the upper normal limit and remained unchanged after TIPS. Baselineportal ET-1 was higher than ET-1 in the peripheral systemic circulation and decreased significantly after TIPS (table 2). In contrast, portal and systemic activation of the renin-angiotensin system (estimated by PRA and AR) were similar and both decreased from an eightfold elevation above normal to a 2–3-fold elevation two

Table 2  Liver and renal function, systemic circulation, and vasoactive mediators during the first weeks after transjugular intrahepatic portosystemic shunt (TIPS) insertion

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Baseline (n=31)</th>
<th>Week 1 (n=30)</th>
<th>Week 2 (n=30)</th>
<th>Week 4 (n=29)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Liver function</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Child-Pugh score</td>
<td>9.5 (1.4)</td>
<td>9.4 (1.8)</td>
<td>9.3 (1.8)</td>
<td>8.8 (1.8)</td>
</tr>
<tr>
<td>Bilirubin (mg/dl)</td>
<td>3.1 (3.3)</td>
<td>4.4 (6.3)</td>
<td>4.1 (7.5)</td>
<td>3.2 (1.9)</td>
</tr>
<tr>
<td>Albumin (g/dl)</td>
<td>2.9 (0.5)</td>
<td>2.0 (0.5)</td>
<td>3.4 (0.6)*</td>
<td>3.3 (0.7)</td>
</tr>
<tr>
<td>Prothrombin time (%)</td>
<td>74 (22)</td>
<td>72 (25)</td>
<td>76 (21)*</td>
<td>77 (20)</td>
</tr>
<tr>
<td>Ascites score</td>
<td>3 (0)</td>
<td>2.7 (0.4)</td>
<td>2.4 (0.6)*</td>
<td>2.1 (0.6)</td>
</tr>
<tr>
<td>Renal function</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Serum creatinine (mg/dl)</td>
<td>2.3 (1.7)</td>
<td>1.7 (1.4)*</td>
<td>1.6 (1.1)*</td>
<td>1.5 (1.2)*</td>
</tr>
<tr>
<td>Creatinine clearance (ml/min)</td>
<td>18 (15)</td>
<td>42 (42)*</td>
<td>84 (48)*</td>
<td>44 (28)*</td>
</tr>
<tr>
<td>Serum urea (mg/dl)</td>
<td>68 (42)</td>
<td>52 (33)*</td>
<td>47 (32)</td>
<td>55 (46)*</td>
</tr>
<tr>
<td>Sodium excretion (mmol/24 h)</td>
<td>12 (16)</td>
<td>38 (29)*</td>
<td>77 (78)</td>
<td>91 (60)*</td>
</tr>
<tr>
<td>Urine volume (ml/24 h)</td>
<td>544 (373)</td>
<td>788 (692)*</td>
<td>1041 (625)*</td>
<td>1248 (621)*</td>
</tr>
<tr>
<td>Systemic circulation</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Heart rate (beats/min)</td>
<td>80.2 (8)</td>
<td>79.7 (10)</td>
<td>84.0 (10)*</td>
<td>82.9 (10)</td>
</tr>
<tr>
<td>Mean arterial pressure (mm Hg)</td>
<td>69.4 (9)</td>
<td>69.4 (7)</td>
<td>75.0 (9)*</td>
<td>74.9 (7)*</td>
</tr>
<tr>
<td>Vasoactive mediators</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ET-1 (pmol/l)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Systemic</td>
<td>1.31 (0.5); n=20</td>
<td>1.39 (0.9); n=20</td>
<td>1.32 (0.4)*; n=20</td>
<td></td>
</tr>
<tr>
<td>Portal</td>
<td>2.05 (0.7); n=20</td>
<td>2.13 (0.8)*; n=20</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PRA (ng/ml/3 h)*</td>
<td>49 (34); n=12</td>
<td>26 (26)*; n=12</td>
<td>26 (26)*; n=12</td>
<td></td>
</tr>
<tr>
<td>Systemic</td>
<td>48 (31); n=12</td>
<td>28 (25)*; n=12</td>
<td>28 (25)*; n=12</td>
<td></td>
</tr>
<tr>
<td>Portal</td>
<td>199 (120); n=8</td>
<td>112 (117)*; n=8</td>
<td>112 (117)*; n=8</td>
<td></td>
</tr>
</tbody>
</table>

Parameters (mean (SD)) of liver and renal function, and systemic circulation (heart rate, mean arterial pressure (MAP)) before and up to four weeks after TIPS. Baseline endotelin-1 (ET-1) and renin-angiotensin system activation (estimated by plasma renin activity (PRA) or active renin (AR)) in the systemic venous and portal circulation were determined in the last consecutive 24 patients and repeated two weeks after TIPS in 20 patients.

Within four weeks, MANOVA detected significant improvements in all renal function parameters, ascites score, and MAP, whereas changes in biochemical liver function parameters showed only a trend towards transient deterioration. Differences over time versus baseline were compared for renal function parameters, ascites score, and MAP for one, two, and four week intervals, whereas all other parameters were tested once for the two week interval.

Differences in liver and renal function v baseline: *p<0.05, **p<0.01, ***p<0.001.

Vasoactive mediators: PRA; baseline v control (n=12); AR; baseline v control (n=8).

Parameters over time versus baseline: §§Portal v systemic ET-1 baseline levels, p<0.01; ¶control v baseline, p<0.05; ¶¶control v baseline, p<0.01.

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SHUNT PATENCY AND ADVERSE EVENTS AFTER TIPS

During the first year of follow up we observed major shunt stenosis in six patients and complete shunt occlusion in one man. TIPS function was successfully re-established by balloon dilatation (three) or by stent prolongation (three) in six patients, while in one patient lethal sepsis excluded reintervention at the time of diagnosis of TIPS dysfunction.

Within four weeks after TIPS, patients experienced on average a mild transient deterioration in some liver function parameters (table 2). During the first three months after TIPS we observed de novo severe encephalopathy in three patients or deterioration in pre-existing mild to moderate encephalopathy in eight patients. Six of these patients had limited short term episodes managed by lactulose but five experienced progressive liver failure. Of note, six of these 11 patients had continued alcohol abuse after TIPS and hospital release.

With respect to non-TIPS patients, renal function worsened progressively in all but one patient. One young man with histological signs of acute alcohol hepatitis and chronic liver cirrhosis experienced gradual recovery from type I to type II HRS. He was the only non-TIPS patient who survived more than three months after HRS onset.

One young man with histological signs of acute alcohol hepatitis and chronic liver cirrhosis experienced gradual recovery from type I to type II HRS. He was the only non-TIPS patient who survived more than three months after HRS onset.
medication with furosemide (40–80 mg/day) and spironolactone (100–200 mg/day) was
weeks after TIPS were determined without diuretics. Thereafter, moderate diuretic

patients with more advanced liver dysfunction at baseline (log rank 18.3; p<0.001) (fig 2A,
2D). After TIPS, three, six, 12, and 18 month survival rates were 81%, 71%, 48%, and 35%
(mean survival 92 (16) weeks; 95% CI 60–123; median 49) compared with non-TIPS patients
(three month survival of 10% and a mean survival of 12 (8.5) weeks (95% CI 0.01–28;
median 2.0)). Type II HRS patients had a better chance of survival than type I patients (log
rank 5.04; p=0.025) (fig 2B). TIPS patients whose ascites were mobilised within one month
(clinical responder) survived longer than patients who did not respond (log rank 46.6;
p<0.001) (fig 2C). Interestingly, even type I patients treated with TIPS (n=14) achieved
three, six, and 12 month survival rates of 64%, 50%, and 20%, which were significantly better
than the survival rates of type I patients who did not undergo TIPS (n=7; p<0.01) with
similar renal dysfunction at baseline (fig 2D; direct Kaplan-Meier comparison not plotted).
Combined survival analysis of all 21 type I HRS patients showed three, six, and 12 month
survival rates of 48%, 38%, and 16% (Kaplan-Meier plot not shown).

For the TIPS patients, multivariate Cox regression analysis, including baseline factors
that were univariately related to death (bilirubin, prothrombin time, Child-Pugh, type of
HRS), revealed only bilirubin (p<0.001) and type of HRS (p<0.05) as independent survival
predictors. Most deaths after TIPS (83%) were related to progressive liver failure. One patient
suffered a procedure related fatal liver haemorhage and two TIPS patients died from
non-liver related events (week 22, cancer progress; week 45, sepsis caused by colon perforation as
diagnosed at autopsy). Subgroup survival analysis in TIPS patients showed that
three month survivors had better baseline liver function (bilirubin 1.7 (1) v 8.7 (5.3) mg/dl,
p<0.001; Child-Pugh score 9.2 (1.3) v 10.8 (1), p=0.08) but similar renal dysfunction
(creatinine clearance 20 (12) v 18 (15) ml/min, NS) compared with non-survivors.

SURVIVAL ANALYSIS
Survival rates (Kaplan-Meier) of the total cohort were 63%, 56%, 39%, and 29% after
three, six, 12, and 18 months, respectively, with a mean survival of 75 (14) weeks (95% CI
48–102, median 34). TIPS patients showed significantly better survival than non-TIPS
weeks after TIPS, despite reintroducing diuretic therapy (table 2).

Discussion
In this phase II study, we have reported on a larger cohort of non-transplantable cirrhotics
with HRS diagnosed according to recent consensus criteria. The median follow up of
two years after HRS diagnosis provides the

Figure 1 Mean (SD) serum creatinine levels (mg/dl; to convert values to µmol/l multiply by 88.4), creatinine clearance (ml/min), and sodium excretion (mmol/24 hours) up to one
year after diagnosis of hepatorenal syndrome in 41 patients of whom 31 received a
transjugular intrahepatic portosystemic stent-shunt (TIPS) and 10 were excluded from
treatment. Six of these 10 persistent alcoholics still abused alcohol after successful TIPS
placement. For the TIPS patients, multivariate Cox regression analysis, including baseline factors
that were univariately related to death (bilirubin, prothrombin time, Child-Pugh, type of
HRS), revealed only bilirubin (p<0.001) and type of HRS (p<0.05) as independent survival
predictors. Most deaths after TIPS (83%) were related to progressive liver failure. One patient
suffered a procedure related fatal liver haemorrhage and two TIPS patients died from
non-liver related events (week 22, cancer progress; week 45, sepsis caused by colon perforation as
diagnosed at autopsy). Subgroup survival analysis in TIPS patients showed that
three month survivors had better baseline liver function (bilirubin 1.7 (1) v 8.7 (5.3) mg/dl,
p<0.001; Child-Pugh score 9.2 (1.3) v 10.8 (1), p=0.08) but similar renal dysfunction
(creatinine clearance 20 (12) v 18 (15) ml/min, NS) compared with non-survivors.
The majority of our cohort experienced sustained improvement in kidney function together with a remarkably high overall one year survival rate of almost 40%. We attribute this benefit to the fact that we considered about 75% of our patients eligible for TIPS as rescue treatment, of whom 77% (24/31) (that is, 59% of the total cohort) exhibited rapid and sustained improvement in GFR and sodium excretion. This response allowed reintroduction of mild diuretic therapy and even withdrawal from haemodialysis in four of seven patients. However, 25% of our high risk patients were excluded from undergoing TIPS because of poor liver function, and almost 25% of TIPS patients did not respond. Both of the latter groups showed a poor prognosis (fig 2C, 2D) similar to previous reports,1 with a median survival of approximately 2 weeks. As our total cohort was comparable with earlier large HRS series3 5 7 with respect to renal and liver function, it is unlikely that the more favourable outcome resulted primarily from patient selection. In addition, even overall survival in the subgroup of our 21 type I HRS patients (14 treated by TIPS) was remarkably high, with three and six month survival rates of 48% and 38% compared with the largest series of HRS patients3 (in which probably most but not all had type I HRS) which showed a three month survival rate of 9% under medical treatment (long term survival rates were not given; median survival 1.7 weeks). Thus it is likely that former HRS series probably included a high proportion of patients who today might benefit from TIPS as a salvage measure.

Although liver transplantation is the treatment of choice for cirrhotic patients with HRS, less invasive approaches have been evaluated because many of these patients are non-transplant candidates (for example due to advanced age, cachexia, and/or active alcoholism). Alternative non-surgical medical approaches using prostaglandins, vasopressin analogues, dopamine, octreotide, or N-acetylcysteine are based mainly on short term applications in small series.2 19–26 Data on vasoactive drugs used as a bridge to transplantation are promising but still unclear.27 28 Long term administration of vasopressin analogues appears to be effective but is often contraindicated or limited by severe vasoconstrictive side effects.20–23 TIPS is a more recent option for the treatment of HRS by semi-invasive portal decompression. Compared with other series, our study provides long term data based on the largest cohort of cirrhotics with HRS diagnosed as type I or type II according to current consensus criteria.2 Of note is the fact that outcome was not influenced by timely transplant rescue within the first critical months after HRS diagnosis, as at baseline all patients were

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**Figure 2.** Kaplan-Meier survival analysis. (A) Cohort of patients with hepatorenal syndrome (HRS) treated by transjugular intrahepatic portosystemic stent-shunt (TIPS) (n=31); (B) survival analysis after TIPS stratified according to HRS subtypes at baseline; (C) survival analysis according to clinical response (improved sodium excretion and ascites control within one month) or no response after TIPS; (D) survival analysis of non-TIPS patients (n=10: type I HRS, n=7; type II HRS, n=3) receiving the best medical support. *p* values were derived from subgroup comparisons using the log rank test.

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TIPS for hepatorenal syndrome
non-transplant candidates. It is well known that portal decompressive surgery (that is, a portacaval end-to-side anastomosis or a side-to-side shunt) not only prevents recurrent bleeding but also prevents the formation of ascites in a larger number of patients than therapies that do not lower portal hypertension. As HRS is the extreme form of refractory ascites, it is sensible to treat HRS with TIPS, especially as this procedure does not carry the risk of open surgery.

Timely TIPS insertion is probably crucial for successful HRS management. Based on our data, recovery was still possible 4–6 weeks even after type I HRS onset in anuric patients who were bridged to TIPS by haemodialysis. However, a clear time limit for TIPS has yet to be defined. As all HRS patients have severe ascites and suffer spontaneously reduced food consumption, chronic fatigue with prolonged bed rest, and progressive muscle atrophy, serum creatinine and urea levels are low in relation to endogenous creatinine clearance data (tables 1, 2). This discrepancy may lead to diagnostic overestimation of GFR with delayed HRS diagnosis.

The pathogenesis of HRS is still incompletely understood. Cirrhotic patients with HRS suffer from peripheral and splanchnic vasodilatation combined with severe portal vasoconstriction mediated by several vasoconstrictive factors such as renin, catecholamines, endothelins, and prostaglandin derivatives. In this situation TIPS insertion not only leads to portal decompression but also to better refilling of the central venous system, although the peripheral hyperdynamic status persists.

Improved refilling can decrease several vasoconstrictors and thus contribute to better renal perfusion leading to an increase in GFR and sodium excretion. These considerations are supported by the observed lesser activation of the renin-angiotensin system after TIPS insertion in our patients despite reintroduction of diuretics, which probably blunts further renin decline to a certain degree. Beneficial short term effects of TIPS on renin activity have been reported in patients with refractory ascites and with HRS. Interestingly, we detected a significant reduction in portal ET-1 levels, while elevated systemic ET-1 levels remained unchanged after TIPS. Similar differences in ET-1 levels between the portal and renal compartment have been described in cirrhotics with ascites after TIPS. Both observations indicate that local or paracrine ET-1 may be more relevant for kidney dysfunction in HRS than circulating ET-1. Furthermore, a decrease in sinusoidal pressure after TIPS may reduce renal sodium retention of the proximal tubule mediated by a sympathetic nervous pathway. Indeed, animal experiments suggest such a hepatorenal reflex, as an acute rise in sinusoidal pressure caused immediate sodium retention. Also, in humans, an increase in short term portal pressure induced an immediate reduction in renal perfusion. Despite potential benefits in renal function, TIPS affects portal venous liver perfusion and thereby carries the risk of progressive liver failure. Therefore, we restricted its placement to patients whose livers were considered to have enough residual capacity to counterbalance reduced perfusion. We excluded patients with Child-Pugh scores >12, bilirubin levels ≥15 mg/dl, or those with severe spontaneous encephalopathy. We are aware that these criteria have been arbitrarily set and that the bilirubin limit in particular was quite generous but reasonable for the given desperate therapeutic situation. Regarding the Child-Pugh score criteria, another report applied similar restrictions. However, we need more controlled data to improve selection regarding the degree of liver dysfunction. In our series early mortality in patients with high baseline bilirubin levels was still substantial after TIPS. This result is similar to outcome data in other reports on shunting or endoscopic treatments for portal hypertension.

Thus prior to TIPS treatment, bilirubin should preferably be stable below 5 mg/dl. We are aware that this criterion probably excludes many cirrhotic patients with HRS and concomitant acute alcoholic hepatitis from early TIPS. However, these patients may become TIPS candidates when HRS persists even after recovery of liver function under strict alcohol abstinence. Such a consideration is an option for transient dialysis treatment, at least in some patients. In the majority of our stented patients with stable bilirubin levels prior to TIPS we observed only mild and transient deteriorations in liver function applying small diameter shunts with limited initial portal decompression (35–45%, measured for technical reasons as reduction in portal vein to vena cava superior pressure gradient). Compared with initial mean portal pressure gradient reductions of approximately 50–60% reported in other TIPS series, our more cautious portal decompression together with our selection criteria for TIPS treatment may have contributed to the rather favourable long term outcome of our total HRS cohort.

In conclusion, limited portal decompression by TIPS represents a promising new option for treating selected HRS patients with sustained efficacy, even without liver transplantation, and warrants further prospective controlled trials in non-transplant and even in transplant candidates. We believe that our long term data provide an adequate basis to plan such controlled trials on the role of TIPS in the management of HRS.

The results of this work were presented in part at the 47th Annual Meeting of the American Association for the Study of Liver Diseases held in Chicago (Hepatology 1996;24:A1292) and preliminary six month renal function data appeared as a research letter in The Lancet (1997;349:697–8).

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3 Gonwa TA, Morris CA, Goldstein RM, Husberg BS, Klintmalm GB. Long-term survival and renal function following
TIPS for hepatorenal syndrome


Long term outcome after transjugular intrahepatic portosystemic stent-shunt in non-transplant cirrhotics with hepatorenal syndrome: a phase II study

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