ABSTRACT

In advanced cirrhosis, increased levels of vasodilators and impaired cardiac compensatory response decrease effective arterial blood volume, causing vasoconstriction of renal arteries and kidney failure in up to 40% of patients after 5 years of follow-up. Hepatorenal syndrome (HRS) diagnostic criteria are: cirrhosis with ascites; serum creatinine (SCr) > 1.5 mg/dL (with no improvement 2 days after diuretic withdrawal and albumin administration). Shock, nephrotoxics and acute parenchymal kidney disease must be excluded. The HRS is classified in: type 1, defined by a 100% increase in SCr to > 2.5 mg/dL in < 2 weeks, and type 2, with a slower and milder decrease in kidney function. Type 3 HRS is an emerging concept, referring to HRS in patients with coexistent kidney disease. Left untreated, average survival of type 1 HRS is 2 weeks whereas in type 2 it is 6 months. Treatment of HRS lies on reversal of the hepatic disease or liver transplantation (combined liver-kidney transplant may be appropriate for patients who have been on renal replacement therapy (RRT) for more than 8 weeks). However, with today’s available therapy, there may be reversibility of HRS without liver transplant. Type 1 HRS is treated with vasoconstrictors (mainly terlipressin; noradrenalin may be an alternative in patients in intensive care units) and albumin. Reversal of HRS occurs in about half of patients. If SCr does not decrease and patients have classic indications for dialysis, RRT can be used as a second-line treatment until liver recovery or transplant. MARS (molecular adsorbent recirculating systems) and Prometheus systems should be considered experimental. Type 2 HRS treatment is based on repeated large-volume paracentesis and albumin administration. If ineffective, vasoconstrictors are used. Since renal impairment is mild, RRT is not indicated. If liver recovery/transplant are unfeasible, patient’s treatment should avoid futilities.

Key words: Dialysis; hepatorenal syndrome; kidney failure; liver cirrhosis; terlipressin.
INTRODUCTION

In advanced cirrhosis, portal hypertension causes severe vasodilation of the splanchnic arteries, leading to a decrease in effective arterial blood volume and arterial pressure. This leads to an intense stimulation of the renin-angiotensin and sympathetic nervous systems, which cause vasoconstriction of the renal arteries and kidney failure. The high levels of plasma renin activity, plasma aldosterone concentration and plasma norepinephrine would also be expected to cause a hyperdynamic circulation, with an increase in heart rate, ventricular contractility, and cardiac output in order to compensate for hypotension. However, whereas cardiac output can increase in early stages of cirrhosis, studies in HRS patients show no increase in heart rate and, actually, a decrease in cardiac output. Therefore, three key mechanisms seem to contribute to HRS: splanchnic vasodilation with hypotension and reduced renal perfusion, renal artery vasoconstriction and cardiac inability to compensate, so that kidney failure seems to result from haemodynamic imbalance, with a preserved tubular function. The kidney failure in HRS is considered a functional defect because there is reversibility of the condition with vasoconstriction of the splanchnic circulation or with liver transplant. Although there is a common misconception that the kidneys are histologically normal, a relatively specific but subtle and reversible renal lesion has been described—glomerular tubular reflux.

In advanced cirrhosis, the vasodilation of the splanchnic arteries is caused by: 1) greater production and activity of vasodilators such as (the most important) nitric oxide and others, such as endogenous cannabinoids and carbon monoxide; 2) proinflammatory cytokines with vasodilatation activity produced in response to bacterial translocation from the intestinal lumen to mesenteric lymph nodes; 3) neoangiogenesis in mesenteric arteries and impaired response to vasoconstrictors. The vasodilators spread along the systemic circulation leading to a global decrease in systemic vascular resistance. These mechanisms constitute the “Classical Peripheral Arterial Vasodilation Hypothesis”.

As stated earlier, impaired inotropic and chronotropic cardiac responses are also important, and led to a revision of this traditional hypothesis. The pathogenesis of the impaired cardiac response in HRS is largely unknown. Contributing factors to reduced
cardiac output may be: i) organic – attenuated systolic and diastolic responses to stress stimuli resulting from the cirrhotic cardiomyopathy, common in patients with HRS; ii) functional – related to a decrease in venous return. Supporting this theory are the facts that: i) HRS occurs in the setting of a decrease in cardiopulmonary pressures, which is compatible with a fall in cardiac preload; ii) intravenous albumin associated with vasoconstrictors and TIPS are included in the treatment of HRS, and both these strategies increase venous return. The impairment in chronotropic cardiac function is probably related to a down regulation of β-adrenergic receptors secondary to the chronic stimulation of the sympathetic nervous system3.

## EPIDEMIOLOGY

The incidence of HRS is highly variable depending on the studies (10 to 40% after 5 years of follow up of a population of patients with cirrhosis and ascites). In a prospective study of 229 nonazotemic patients with cirrhosis and ascites the hepatorenal syndrome developed in 18 and 39 percent at one and five years, respectively12. Patients with hyponatremia and a high plasma renin activity presumably reflected a neurohumoral activation presumably reflected a more severe decline in effective perfusion13,14.

The hepatorenal syndrome characteristically occurs in patients with advanced hepatic disease and in the presence of portal hypertension (hepatic cirrhosis) and / or hepatic insufficiency (severe alcoholic hepatitis, hepatic metastases or fulminant hepatitis from any cause)13,15,16. Although hepatorenal syndrome can be seen in most forms of severe hepatic disease, patients with primary biliary cirrhosis appear relatively protected17, possibly due in part to the natriuretic and renal vasodilator actions of retained bile salts.

## DIAGNOSTIC CRITERIA

To be diagnosed with HRS, patients have to fulfill all the criteria18 presented in Table I, as proposed by the International Ascites Club in 2007. When comparing these criteria with the former ones (published in 1996)5, there was a significant advance in terms of simplification of the diagnosis, hence permitting an earlier, more effective treatment. The more important differences between the old and the new criteria are described below. In the 1996’s criteria, there had to be a serum creatinine of > 1.5 mg/dL (≥ 133 μmol/L) after at least 2 days with diuretic withdrawal and volume expansion with albumin. The recommended dose of albumin is a single infusion of 1 g/kg of body weight (maximum, 100 g).

### Table I

<table>
<thead>
<tr>
<th>Hepatorenal Syndrome – diagnostic criteria3:</th>
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<tbody>
<tr>
<td>1. Cirrhosis with ascites</td>
</tr>
<tr>
<td>2. Serum creatinine &gt; 1.5 mg/dL (≥ 133 μmol/L)</td>
</tr>
<tr>
<td>3. No improvement in serum creatinine level (decrease to ≤ 1.5 mg/dL ≤ 133 μmol/L) after at least 2 days with diuretic withdrawal and volume expansion with albumin. The recommended dose of albumin is a single infusion of 1 g/kg of body weight (maximum, 100 g)</td>
</tr>
<tr>
<td>4. Absence of shock</td>
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<tr>
<td>5. No current or recent treatment with nephrotoxic drugs</td>
</tr>
<tr>
<td>6. Absence of parenchymal kidney disease as indicated by proteinuria &gt; 500 mg/d, microhaematuria (&gt; 50 red blood cells/high-power field), and/or abnormal renal ultrasonography. However, the urine sediment may show a variety of abnormalities, such as haematuria due to bladder instrumentation and underlying coagulopathy (bladder catheters are necessary only when there is marked oliguria), and granular casts due to hyperbilirubinemia, so an accurate clinical judgment is warranted. Also, hepatorenal syndrome can occur in patients with coexisting chronic kidney disease (see type 3 Hepatorenal Syndrome).</td>
</tr>
</tbody>
</table>

The 1996’s criteria specifically referred that was necessary to exclude gastrointestinal or renal losses of fluid whereas in the new diagnostic criteria the clinician focuses on whether volume expansion improves creatinine. Thirdly, in the 1996’s diagnostic criteria, an acute infection actually excluded HRS. Nowadays we know that only sepsis must be excluded, as the most common trigger for the development of type 1 HRS is actually bacterial infection, particularly spontaneous bacterial peritonitis (SBP). This way, treatment can (should) be started without waiting for complete recovery from the infection. Finally, in the 1996’s diagnostic criteria, existed additional criteria (low urinary volume, serum and urinary sodium – U Na < 10 mEq/day – and high urine to plasma osmolality ratio).
Although often present, these parameters are not essential for the diagnosis and so were removed. Looking again at Table I, we realize the efforts made to simplify the diagnostic criteria and start treatment as soon as possible. However, HRS remains a diagnostic of exclusion and a diagnostic marker is still lacking.

**CLINICAL TYPES OF HRS**

**Type 1**

Rapidly progressive decrease in kidney function: 100% increase in serum creatinine to a final value > 2.5 mg/dL (> 221 μmol/L) in < 2 weeks. The clinical presentation is usually that of acute kidney failure.

**Type 2**

Stable or slowly progressive decrease in kidney function that does not meet the criteria of type 1. The clinical picture is that of ascites refractory to diuretic therapy.

**Type 3**

Some authors have singled out a type 3 hepatorenal syndrome, in which there is coexistent kidney disease and hepatorenal syndrome. Although at first sight this may seem odd (because in the diagnostic criteria one has to exclude the presence of parenchymal renal disease), one has to consider that, for example, diabetics with diabetic nephropathy and non-alcoholic fatty liver disease may develop hepatorenal syndrome. Other systemic diseases may similarly affect both the liver and the kidney. In fact, a recent study found that 85% of end-stage cirrhots had pre-existing intrinsic renal disease on renal biopsy. In some cases renal biopsy may be necessary for diagnosis and for selection of patients for combined liver-kidney transplant.

**DIFFERENTIAL DIAGNOSIS (Table II)**

The hepatorenal syndrome is a diagnosis of exclusion, and other diseases need to be considered, namely:

- Severe sepsis
- Other causes of severe kidney failure that can arise in patients with advanced cirrhosis:
  - Drug-induced nephrotoxicity (NSAIDs, aminoglycosides, radiological contrasts, others).
  - Pre-renal failure due to volume depletion (from diarrhoea, vomiting, increased diuresis due to use of diuretics or not; other causes of hypovolaemia).
  - Acute Tubular Necrosis (ATN): ATN is usually suspected from the history and from the often rapid rise in the serum creatinine, which contrasts to the usually gradual rise in hepatorenal syndrome. Some of the traditional laboratory methods used to distinguish pre-renal disease from ATN (such as the urinalysis or the fractional excretion of sodium) may not be helpful in patients with hepatic disease.
  - Glomerulonephritis in patients with hepatitis B or C; Immunoglobulin A nephropathy (mainly combined with alcoholic cirrhosis); others

Distinguishing the hepatorenal syndrome from these other disorders is clinically important because of the marked difference in prognosis and the urgent need to begin treatment directed to hepatorenal syndrome.

**CLINICAL PRESENTATION**

**Type 1 HRS**

- Severe and progressive kidney failure. However, due to the marked reduction in creatinine production among such patients, the serum creatinine may increase by as little as 0.1 mg/dL (9 micromol/L) per day, with intermittent periods of stabilization or even slight improvement.
• Severe circulatory dysfunction (mean arterial pressure usually is 70 mm Hg) and very low systemic vascular resistance.
• Severe liver disease, with jaundice, coagulopathy, low albumin levels, hepatic encephalopathy, poor nutritional status, and large ascites and oedema.
• Urine volume usually is not extremely reduced and some patients may have normal urine volumes, with markedly lower output being observed only within a few days from death.25,26

Type 2 HRS:
• Moderately severe kidney failure (serum creatinine levels of ~ 2.0 mg/dL) of functional origin that remains stable for variable periods.
• Ascites, usually resistant to diuretic therapy (because of the combined influence of profound sodium retention, reduced GFR, and markedly increased levels of aldosterone and norepinephrine).
• Dilutional hyponatremia.
• Some patients with type 2 HRS develop type 1 HRS, which may arise spontaneously or as a result of some complication, usually a bacterial infection.

Type 3 HRS:
• Development of HRS in a patient with coexistent kidney disease.

ADDITIONAL DIAGNOSTIC WORKUP
Liver disease may be associated with near normal values for both the BUN (due to decreased urea production) and the serum creatinine (due to muscle wasting) despite a relatively large reduction in GFR27-28. The presence of kidney disease in this setting can be documented by a reduction in creatinine clearance, but significant overestimation of GFR can still occur28-30. Because of the problems with changes in creatinine production and secretion, other endogenous compounds have been evaluated in an effort to provide a more accurate estimation of GFR, including cystatin C and urinary biomarkers such as neutrophil gelatinase-associated lipocalin (NGAL).

Cystatin C: It has been proposed that cystatin C-based equations would be more accurate in patients with cirrhosis when compared with creatinine clearance. For example, in a study in liver cirrhotic patients32, a cystatin C-based formula (developed by Larsson and Hoek) showed significantly lower bias and higher precision than the creatinine-based formula (Cockroft & Gault or MDRD) for GFR estimation. However, both creatinine and Cystatin C-based equations overestimated the true GFR by 105-154%. Therefore, even in populations in which cystatin C would be expected to outperform creatinine based GFR calculations, cystatin C-based equations are not totally accurate.33 Also, even if cystatin C proved to be more accurate for the assessment of GFR than serum creatinine in cirrhotics, whether measurement of cystatin C levels would improve patient care is at present unknown, so cystatin C is not at the moment routinely performed in these patients.

Urinary biomarkers, such as neutrophil gelatinase-associated lipocalin (NGAL), tend to be lower in pre-renal azotaemia and hepatorenal syndrome than in acute tubular necrosis (ATN), but there is considerable overlap between these conditions.34,35

PRECIPITATING FACTORS:
• None (in some patients)
• After effective arterial blood volume is decreased by:
  • Bacterial infections and, in particular, spontaneous bacterial peritonitis. Approximately one-third of patients with spontaneous bacterial peritonitis develop HRS and are treated simultaneously for both disorders. Of these patients, about one-third experience reversal of HRS when the infection is resolved. However, the remainder develop either stable (type 2) or progressive HRS (type 1). Patients who develop type 1 HRS as a result of spontaneous bacterial peritonitis have a dismal outcome, with almost 100% hospital mortality if not treated appropriately. Infections other than spontaneous bacterial peritonitis also may cause HRS, but its frequency and severity usually are lower than that of patients with spontaneous bacterial peritonitis.
• Gastrointestinal bleeding. The development of kidney failure after gastrointestinal bleeding is not very common in patients who have cirrhosis (10%) and it is almost fully confined to patients with hypovolaemic shock. In most instances, it is associated with ischaemic hepatitis, which implies that the kidney failure most likely is related to ATN and not HRS.

• Large volume paracentesis (> 5 L) in the absence of albumin administration. Large-volume paracentesis without albumin may trigger HRS in 15% or more of cases.

Although diuretics have often been mentioned as precipitants of HRS, diuretics do not cause hepatorenal syndrome. Diuretics can, however, cause azotaemia, which improves with the cessation of therapy and fluid repletion, while in the hepatorenal syndrome kidney function typically worsens inexorably, even after diuretics are stopped.

## TREATMENT OF HRS

### Type 1 HRS

Treatment of type 1 HRS can be divided into three stages, depending on the severity of the hepatic and renal disease. These stages are: treatment of the hepatorenal syndrome itself, treatment of the acute kidney injury and treatment of the advanced hepatic disease. The first stage, treatment of the hepatorenal syndrome itself, includes pharmacological treatment (with intravenous albumin, to expand intravascular volume, together with vasoconstrictors, to reverse splanchnic arterial dilatation) and interventional procedures (transjugular intrahepatic portosystemic shunt – TIPS – and peritoneovenous shunt). The second stage, treatment of the acute kidney injury (with renal replacement therapy) is necessary when the pharmacological treatment is ineffective or is still underway in patients who are candidates to kidney transplant or in whom reversal of the hepatic disease is a possibility. The final stage, treatment of the advanced hepatic disease, either by resolving the hepatic insult or by liver transplant, is the only definitive treatment (artificial support of the hepatic and renal function may be temporarily required, in which case the MARS and Prometheus techniques may be of use).

### Treatment of the hepatorenal syndrome itself

#### Pharmacological treatment

**Albumin and vasoconstrictors**

Intravenous albumin in association with vasoconstrictors (Table III) improves survival and is currently considered the best therapy for type 1 HRS. This approach is intended to expand intravascular volume and to cause vasoconstriction of the greatly dilated splanchnic arterial bed. This in turn alleviates arterial underfilling, lessens the activation of the endogenous vasoconstrictor systems, and increases kidney perfusion and GFR.

- Administration of albumin: 1 g/kg body weight at day 1 and 2 – maximum 100g – followed by 25-50 g/d.
- Administration of vasoconstrictor drugs:
  - Terlipressin (vasopressin analogue, acts on V1 vasopressin receptors in vascular smooth muscle cells):
    - 1 mg/4-6 h as IV bolus;
    - The dose is increased up to a maximum of 2 mg/4-6 h after 3 days if there is no response to therapy, defined by a decrease in serum creatinine > 25% of pre-treatment values.
    - Response to therapy is indicated by a marked decrease in the high serum creatinine levels, to less than 1.5 mg/dL (< 133 μmol/L).

#### Table III

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dosage</th>
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<tbody>
<tr>
<td>Albumin</td>
<td>1 g/kg (at day 1 and 2 – max. 100g) followed by 25-50 g/d.</td>
</tr>
<tr>
<td>Terlipressin</td>
<td>1 mg every 4 to 6 h (IV bolus); if after 3 days there is no response, increase the dose up to 2 mg / 4-6h. Continue treatment for 5-15 days (longer treatments may be considered in patients with partial improvement of renal function).</td>
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<tr>
<td>Norepinephrine</td>
<td>0.5-3 mg/h (IV infusion); titrate to increase mean arterial pressure by 10 mmHg. Maintain treatment until serum creatinine decreases to &lt; 1.5 mg/dL (&lt; 133 μmol/L).</td>
</tr>
<tr>
<td>Midodrine + octreotide</td>
<td>Midodrine: 7.5 mg (orally) every 8 hours; dose may be increased up to 15mg 3id. Octreotide: 50 mcg/h (IV infusion); alternatively, 100-200μg 3id (subcutaneously).</td>
</tr>
</tbody>
</table>
- Treatment is usually given from 5-15 days (occasionally longer if there is some but not complete improvement of renal function after two weeks).

- Noradrenaline or midodrine (α-adrenergic agonists, act on α1-adrenergic receptors in vascular smooth muscle cells):
  - Norepinephrine (given in an ICU): 0.5-3 mg/h as continuous intravenous infusion aimed at increasing mean arterial pressure by 10 mmHg. Treatment is maintained until serum creatinine decreases to < 1.5 mg/dL (< 133 μmol/L).
  - Midodrine (a systemic vasoconstrictor): 7.5 mg orally at eight-hour intervals, increased to a maximum of 15 mg 3id if needed.

- Noradrenaline or midodrine (α-adrenergic agonists, act on α1-adrenergic receptors in vascular smooth muscle cells):
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  - Midodrine (a systemic vasoconstrictor): 7.5 mg orally at eight-hour intervals, increased to a maximum of 15 mg 3id if needed.

Treatment with the combination of terlipressin and albumin is associated with reduced mortality and reversal of HRS in 40%-50% of patients, making this approach the preferred initial therapy (according to the European Association for the Study of the Liver – EASL – 2010 guidelines). However, given the fact that some studies found no difference in efficacy and safety between patients treated with terlipressin plus albumin versus noradrenaline plus albumin, and because terlipressin may be significantly more expensive than noradrenaline, noradrenaline is recommended by some authors as first line therapy for patients with HRS who are in a ICU (noradrenalin perfusion is not usually available on the general medical ward). However, in Portugal, this price difference may not be always present so an individual approach is warranted. Response to treatment with terlipressin and albumin is associated with a progressive decrease in serum creatinine concentration, increased urine output, and improvement of hyponatremia. Factors that predict a response to treatment are an increase in arterial pressure during treatment and low baseline creatinine level. After withdrawal of therapy, HRS recurs in < 15% of patients, and in these cases, a second treatment with terlipressin is usually effective. The incidence of side effects (usually ischaemic) that mandate discontinuation of treatment is ~12%. Terlipressin has been associated with an increase of cardiovascular adverse events.

- Terlipressin: 0.5-3 mcg/kg/h as continuous intravenous infusion aimed at increasing mean arterial pressure by 10 mmHg. Treatment is maintained until serum creatinine decreases to < 1.5 mg/dL (< 133 μmol/L).

- Midodrine (a systemic vasoconstrictor): 7.5 mg orally at eight-hour intervals, increased to a maximum of 15 mg 3id if needed.

Alternative therapies

Acutely lowering renal sympathetic tone and renal vascular resistance in the early stages of hepatorenal syndrome by the intravenous administration of the sympatholytic agent, clonidine, can raise the GFR by as much as 25 percent. However, this benefit does not appear to be sustained with chronic oral therapy, despite a persistent reduction in sympathetic activity.

Interventional procedures

TIPS

A recent study showed that vasoconstrictor therapy followed by stent placement (TIPS – transjugular intrahepatic portosystemic shunt) was effective in a limited number of patients with type 1 HRS and type 2 HRS. This approach is sometimes successful in highly selected patients, who fail to respond to vasoconstrictor therapy. However, this procedure is associated with numerous complications (high
incidence of encephalopathy, among others) and, because of the need for intravenous contrast, it may cause acute kidney injury. For this reason, some experts prefer dialysis as a first option (continuous renal replacement therapy) for patients whose serum creatinine remains above 1.5 mg/dL despite medical therapy. Overall, the available results suggest that TIPS should be considered only as a last resort in selected patients. More studies are required to establish the value of TIPS placement in the treatment of HRS.

Peritoneovenous shunt

Peritoneovenous shunts drain peritoneal fluid from the peritoneum into the internal jugular vein, reinfusing ascites into the vascular space. It is now rarely used because of an appreciable rate of complications and lack of evidence that peritoneovenous shunting prolongs patient survival.

Goal of therapy

The goal of medical therapy or TIPS in patients with hepatorenal syndrome is reversal of the acute kidney injury (decrease in the high serum creatinine levels, to least < 1.5 mg/dL, < 133 μmol/L). In addition, when patients are treated with norepinephrine, terlipressin, or midodrine plus octreotide, an immediate goal of therapy is to raise the mean arterial pressure by approximately 10 to 15 mmHg. The magnitude of the increase in mean arterial pressure induced by these vasoconstrictors appears to be significantly associated with the magnitude of the decrease in serum creatinine. As an example, in a systematic review of 501 patients with hepatorenal syndrome from 21 studies, a 9 mmHg increase in mean arterial pressure predicted a 1 mg/dL (88.4 micromol/L) decrease in serum creatinine. If a patient has no improvement in renal function after two weeks, therapy with these drugs can be considered futile.

Treatment of the acute kidney injury – when vasoconstrictors are ineffective

Renal replacement therapy

RRT is not considered the first-line treatment for patients with type 1 HRS because it does not correct the underlying pathogenesis. RRT should be started when patients with type 1 HRS are unresponsive to vasoconstrictors and when there are signs of uremia, volume overload, severe metabolic acidosis, or hyperkalemia. However, not all patients are candidates for dialysis. Dialysis is useful as a bridge to liver transplantation or until there is liver recovery. Bridging patients to liver transplantation includes patients either waiting for a transplant or being evaluated for liver transplantation. In patients who develop a need for RRT but who are not expected to recover liver function / to receive a liver transplant, long-term RRT is usually not indicated / should be withheld except as a trial to see if renal function will return.

Haemodialysis is frequently difficult to perform in patients with hepatorenal syndrome since decompensated hepatic function is associated with haemodynamic instability, thrombocytopenia and coagulopathy. Survival with RRT is poor with only 30–60% of patients surviving to liver transplant. Some success has been accomplished with continuous renal replacement (CRRT) modalities, which have potential advantages such as improved cardiovascular stability, more gradual correction of hyponatraemia (necessary to avoid central pontine myelinolysis), less fluctuation in intracranial pressure and removal of inflammatory cytokines, such as TNF-α and IL-6. Studies, however, do not show superiority of CRRT when compared with conventional intermittent RRT. However, all trials available are non-randomized with populations considered to be non-comparable to one another. Therefore, the decision of which modality to choose continues to be based on the clinical characteristics of the patient as dictated by haemodynamic stability and severity of illness.

Treatment of the advanced hepatic disease – looking for a definitive treatment

Although with today’s available therapy there may be reversibility of HRS without liver transplant, the definitive treatment of hepatorenal syndrome is improvement of liver function (for example, by recovery of alcoholic hepatitis, treatment of decompensated hepatitis B with effective antiviral therapy or recovery from acute hepatic failure), or liver transplantation. In patients in which recovery of liver function may be expected or who are candidates for liver transplantation, temporary artificial support of
the liver (and renal) function may be considered, using either the MARS (molecular adsorbent recirculating system) or FPSA (fractionated plasma separation and adsorption – the Prometheus system).

**MARS and Prometheus**

The MARS system is designed to remove albumin-bound toxins (including vasodilators) by albumin dialysis as well as providing standard continuous renal replacement therapy (CRRT). The albumin dialysate is then regenerated utilizing an anion exchange resin and active charcoal adsorption. MARS has been utilized in treatment of HRS and was shown to be superior to CRRT in terms of patient survival, improved haemodynamics and urine output. However, no large-scale trial has been carried out and more recent studies (6 patients) in patients with type 1 HRS not responding to vasoconstrictor therapy found no improvement following MARS therapy in terms of systemic haemodynamics. Also, in a French study with thirty-two patients with type 1 hepatorenal syndrome, MARS therapy improved renal function in only very few patients with type 1 HRS.

Fractionated plasma separation and adsorption (FPSA) is a method of albumin dialysis that is integrated into an extracorporeal liver support device (Prometheus). In the HELIOS trial, a randomized-controlled European multicenter trial of FPSA therapy, a total of 145 patients with acute-on-chronic liver failure were either treated with standard medical treatment and FPSA eight to 11 times over 21 days or with standard medical treatment alone. There was no statistically significant difference in the overall survival. However, significant survival benefit was observed under FPSA therapy in a predefined subgroup of patients with type 1 hepatorenal syndrome.

In conclusion, although MARS and Prometheus systems may be used to bridge patients to liver transplant, controlled studies are needed. Until then, these therapies should be considered experimental.

**Liver transplant**

Liver transplant is the first choice of treatment for patients with cirrhosis and type 1 HRS because of their low survival expectancy. Therefore, patients who are candidates for liver transplant should be referred immediately to transplant centres.

Because kidney failure is reversible after liver transplant, combined liver-kidney transplant is generally considered appropriate only for patients who have been on RRT for more than 8 weeks who have a low likelihood of recovery of kidney function. However, one must consider that the exact duration of pre-liver transplant kidney dysfunction or dialysis that is amenable to recovery is not known. Retrospective studies from single centres have shown the importance of the duration of > 12 weeks of SCR ≥ 1.5 mg/dL and dialysis ≤ 4 weeks pre-transplant on post-transplant renal outcomes. The duration of pre-transplant dialysis may be variable according to the physician/centre considered, so these results must be viewed with caution. The decision for combined liver-kidney versus liver transplant alone should be undertaken with consideration of duration of HRS, AKI and CKD and risk factors for progression of CKD present at the time of liver transplant such as hypertension, diabetes and obesity. In summary, the 8th international consensus conference of the Acute Dialysis Quality Initiative (ADQI) Group suggest liver transplantation alone for candidates with type-1 HRS for less than four weeks and simultaneous liver-kidney (SLK) for those at risk for non-recovery of renal function (2D).

 Provision of intra-operative continuous RRT during liver transplant may be indicated to help control volume and electrolytes. Use of vasoconstrictors before liver transplant with the aim of performing transplant on patients with normal or near-normal kidney function remains an open question, because studies are scarce and include a small number of patients. However, excellent survival has been reported with the 2 approaches (“transplant-without treating HRS” or “treat HRS before transplant”).

Patients who are not candidates for transplant or who have important comorbid conditions

Decisions about the management of patients who are not candidates for transplant or who have important comorbid conditions should be made on an individual basis. In these patients, therapy with vasoconstrictors should be individualized and RRT probably be reserved for particular cases (potentially...
reversible chronic liver diseases – alcoholic hepatitis, acute-on-chronic liver failure, etc. – with no important associated comorbid conditions) in order to avoid futilities.

**Type 2 HRS**

– Usually managed as outpatients
– Spironolactone and other potassium-sparing diuretics should generally be avoided because of the risk of hyperkalaemia, whereas loop diuretics, such as furosemide, usually lack efficacy. However, diuretics can be given to patients without adverse reactions to diuretics who have a sodium excretion under diuretic treatment of > 30 mEq/d².
– Treatment of ascites is based on repeated large-volume paracentesis and albumin administration (8 g/L of ascites removed)³⁶⁻⁵⁹.
– More studies are required to more fully understand the role that vasoconstrictors plus albumin and TIPS may have in treating type 2 HRS. Some algorithms propose the use of vasoconstrictors in patients with type 2 HRS who are candidates for liver transplant and in whom there is a rise in serum creatinine².
– Renal replacement therapy is not indicated in the management of patients with type 2 HRS because of the lack of a severe decrease in kidney function.

**Prevention of HRS²,²⁴,³⁶:**

– Administration of IV albumin to all patients with cirrhosis and spontaneous bacterial peritonitis (1.5 g/kg body weight at diagnosis and 1 g/kg 48 hours later) – as proposed in the guidelines of the European Association for the Study of the Liver (EASL) – reduces kidney impairment and improves survival.
– Long-term oral administration of norfloxacin (400 mg/d) in patients with ascitic fluid protein < 15 g/L and associated decreased liver and/or kidney function (bilirubin > 3 mg/dL [> 51.3 μmol/L], Child-Pugh score > 10, serum sodium < 130 mEq/L [< 130 mmol/L], and/or serum creatinine > 1.2 mg/dL [> 106.1 μmol/L]) reduces the risk of HRS and improves survival. These effects probably are related to prevention of bacterial translocation, suppression of pro-inflammatory cytokines, and improvement in circulatory function.
– One report suggested that the reduction in intrahepatic pressure induced by transjugular intrahepatic portosystemic shunt (TIPS) placement may prevent the development of the hepatorenal syndrome. This retrospective study evaluated 204 patients with variceal bleeding who were treated with either a portosystemic shunt or sclerotherapy (or other non-shunt modalities)⁶⁰. Portasystemic shunting was associated with a lower incidence of ascites (15 versus 73%) and hepatorenal syndrome (4 versus 21%), a higher incidence of encephalopathy, and no difference in overall patient survival⁶⁰.

**Prognosis**

– If untreated, median survival of type 1 HRS is only 2 weeks. In type 2, average median survival is 6 months. The best hope for reversal of the renal failure is an improvement in hepatic function due to partial resolution of the primary disease or to successful liver transplantation. The rate of recovery of kidney function following recovery of liver failure is uncertain. However, a substantial proportion of patients who have progressed to dialysis and survive to receive a liver transplant do recover kidney function⁶¹.

**Conflict of interest statement:** None declared.

**References**

What’s new in hepatorenal syndrome? An updated review for the nephrologist


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