Introduction

The advent of successful orthotopic liver transplantation for patients with both acute and chronic liver failure has been a major breakthrough in clinical practice, for patients who would have otherwise died [1]. Patients with chronic liver disease are now listed for transplantation well before terminal liver failure, and therefore can potentially wait for a suitable donor liver. However, those with acute hepatic failure [2] can be desperately ill, and may require urgent transplantation. Unfortunately, the number of patients with liver disease awaiting transplantation now exceeds the donation rate. If no liver is immediately available, and the clinical condition of the patient is deteriorating, then the question of artificial liver support arises. Similarly, some patients will have contraindications to transplantation, due to cardiovascular or intracranial instability, sepsis or psychosocial factors [3]. In these cases artificial liver support may help, and so allow subsequent transplantation. Patients with chronic liver disease may develop encephalopathy and hepatorenal syndrome [4] secondary to acute events including sepsis and gastrointestinal hemorrhage. Artificial liver support may help to quicken recovery from such acute events.

Over the years several treatments have been advocated to improve survival in both patients with acute and chronic liver failure. After initial success reported in small selected series, many have not then been substantiated in larger randomized controlled trials.

Dialysis

The introduction of peritoneal and standard intermittent hemodialysis was shown not to affect the outcome in patients with hepatorenal syndrome [5]. Similarly, more recent series have not shown any significant impact on outcome [6]. Peritoneal dialysis has been advocated in patients with liver failure, however there is both an increased risk of peritonitis [6], coupled with inefficient dialysis due to the changes in peritoneal blood flow. Indeed, when two-liter exchanges are performed, patients may exhibit marked cardiovascular instability [7], thus tidal peritoneal dialysis is preferred.

Conventional dialysis preferentially removes small water-soluble solutes, but not those, which are lipid soluble or bound to albumin. Early reports suggested that hemodialysis using polyacrylonitrile dialysers improved conscious level, and that this was due to the removal of middle-sized solutes [8]. Unfortunately these treatments were often complicated by severe hypotension, and many patients died of cerebral edema.

To improve solute removal during dialysis, Mitzner and colleagues [9] in Rostock developed an albumin-based dialysate, molecular adsorbent recycling system (MARS). This system has been shown to remove bilirubin, bile acids, aromatic amino acids and free fatty acids, by transfer from plasma albumin to dialysate albumin, using a high flux polysulphone dialyzer. Thereafter, the effluent dialysate albumin is regenerated by passage...
through a low flux dialyzer against fresh bicarbonate dialysate and then two charcoal columns. A small randomised prospective controlled trial compared additional MARS treatment to hemodiafiltration in a group of patients with severe chronic liver disease. Although mortality was significantly reduced in the MARS-treated group after 7 and 30 days, by 40 days only 1 patient was still alive [9]. Another group just used fresh albumin-based dialysate and treated 3 patients who had decompensated chronic liver disease [10]. In all 3 cases encephalopathy resolved and two were subsequently successfully transplanted. Both systems reported a mild degree of thrombocytopenia associated with treatment.

**Hemofiltration**

Hemofiltration removes a larger spectrum of watersoluble solutes than hemodialysis. Early reports suggested improvement in hepatic encephalopathy and survival in patients with fulminant hepatic failure treated by hemofiltration, and compared to dialysis the nonsurvivors predominantly died of multiorgan failure with severe hepatic necrosis rather than cerebral edema [11]. More recent studies have not borne out the earlier success [12].

Subsequently continuous forms of renal replacement therapy (CRRT) have been developed. Although these resulted in greater cardiovascular and intracranial stability compared to intermittent hemodialysis and/or hemofiltration, they were not shown to significantly improve survival [7]. CRRT did allow stabilisation of the patient, possibly due to thermal losses rather than toxin removal, and allowed the controlled correction of hyponatremia prior to transplantation [7].

**Charcoal**

Charcoal hemoperfusion was introduced to try and remove lipid soluble toxins. Several promising studies were published from the early 1980s up to 1992 in patients with acute liver failure, with reports of reduction in grade of encephalopathy and improved survival. However, when the same center then conducted a prospective randomized controlled trial, they showed that charcoal hemoperfusion did not have any additional benefit over conservative management [13].

The BioLogic-DT sorbent system utilises a cellulosic membrane dialyzer and both a charcoal suspension and a sodium-loaded cation exchange resin as a sorbent dialysate [14]. In theory, only those solutes which can pass through the cellulosic membrane can be removed. Although there are small studies suggesting an improvement in patients with chronic liver disease who have decompensated [14], others in acute liver failure have failed to show any benefit [15].

**Plasma Exchange**

Although high volume plasma exchange and/or plasma immunoabsorption have been reported to improve cardiovascular and intracranial stability in patients with acute liver failure, the results from Copenhagen did not show any significant improvement in patient survival compared to conservative management [16]. Similarly, the results from the Japanese series do not show an obvious improvement with plasma perfusion, apart from a small group with acute viral hepatitis who had 70–80% survival [17].

**Extracorporeal Hepatic Perfusion**

Extracorporeal whole organ perfusion for acute liver failure was first described in 1957. Thereafter, many short uncontrolled series were published using baboon or pig livers, but this technique lost favor due to technical problems. More recently, a case of human extracorporeal perfusion with a human liver that was not suitable for transplantation was used in a young child with acute liver failure [18]. The liver survived for 72 h during which time the child’s intracranial pressure declined, metabolic acidosis corrected, prothrombin time normalised, and coagulation factors V and VII increased. Thus, this treatment may act as a bridge to transplantation.

**Hepatocytes**

Two bioartificial livers have been developed, one using primary porcine hepatocytes, the other a transformed human hepatoma cell line (C3A) [19]. To prevent clotting in the porcine system, blood passes through a plasma filter, and the plasma filtrate then perfuses through the porcine bioartificial liver (BAL), and then through a charcoal adsorption column before returning to the patient [20]. Whereas in the extracorporeal liver assist device (ELAD), blood passes directly through two cartridges of human hepatocytes. The initial pilot studies of these devices
showed that there was some hepatocyte function. However, neither pilot data suggested a major improvement in patient outcome. However, this may have been due to the amount of functioning hepatocyte mass. The initial trials with the ELAD used a single cartridge containing approximately 200 g of hepatocytes, subsequent trials used 400 g and the current planned multicenter trial will use 800 g.

One other treatment described as a potential bridge to transplantation is the injection of human hepatocytes, from cadaveric and human donor livers that are unsuitable for solid organ transplantation. Hepatocytes can be injected percutaneously into the spleen or into the splenic vein, and seed into the liver [21].

### Summary

The liver plays a central role in metabolic homeostasis, but also a major role in host-defence mechanisms. There is great debate as to the underlying etiology of what causes the clinical syndrome of acute liver failure, whether it represents toxin accumulation due to defective hepatocyte function, or is due to endothelial and Kupffer cell mediated inflammation secondary to hepatocyte necrosis with resultant endotoxemia, cytokine activation and recruitment of inflammatory cells. Thus, treatments designed to support the patient with liver failure must not only remove toxins, but also provide adequate hepatic function, until sufficient hepatic regeneration occurs, or transplantation.

The history of treating patients with acute hepatic failure is littered with small uncontrolled series advocating one treatment after another. It is important after the debacle of charcoal perfusion that the newer treatments are carefully evaluated by randomized controlled trials. The BioLogic-DT and MARS systems will remove a varying amount of toxins, but do not provide hepatic support. Bioartificial livers, such as the ELAD and hepatocyte infusions, may prove to be the superior treatment, but the question arises as to how many hepatocytes are required.

### References