Acute liver failure (ALF) is a rare, potentially fatal complication of severe hepatic illness resulting from various causes. In a clinical setting, severe hepatic injury is usually recognised by the appearance of jaundice, encephalopathy and coagulopathy. The central and most important clinical event in ALF is occurrence of hepatic encephalopathy (HE) and cerebral edema which is responsible for most of the fatalities in this serious clinical syndrome. The pathogenesis of encephalopathy and cerebral edema in ALF is unique and multifactorial. Ammonia plays a central role in the pathogenesis. The role of newer ammonia lowering agents is still evolving. Liver transplant is the only effective therapy that has been identified to be of promise in those with poor prognostic factors, whereas in the others, aggressive intensive medical management has been documented to salvage a substantial proportion of patients. A small fraction of patients undergo liver transplant and the remaining are usually treated with medical therapy. Therefore, identification of the complications and causes of death in such patients, and use of appropriate prognostic models to identify those who need liver transplant and those who can be managed with medical treatment is a vital component of therapeutic strategy. In this review, we discuss the various pathogenetic mechanisms and treatment options available. (J CLIN EXP HEPATOL 2015;5:S104–S115)

Acute liver failure (ALF) can be a fatal complication of acute hepatic injury and occurs unpredictably. It is a rare clinical entity marked by the sudden loss of hepatic function and a severe life-threatening course in a person with no prior history of liver disease. ALF represents a syndrome rather than a specific disease, having multiple causes that vary in course and outcome. ALF is difficult to identify in its early stages, resulting in frequent delays in initiation of treatment. The causes of ALF include viral hepatitis, drug induced and toxin-induced liver damage, metabolic errors, ischemia, and miscellaneous rare causes.

ALF is defined by three criteria: (1) rapid development of hepatocellular dysfunction (jaundice, coagulopathy), (2) encephalopathy, and (3) absence of a prior history of liver disease. However, the interval between onset of acute hepatic injury (jaundice) and onset of liver failure (encephalopathy with or without coagulopathy) in such patients (icterus-encephalopathy interval; IEI) has been described to be between 4 weeks (Indian definition) to 24 weeks (AASLD–ALF study group). Further, due to the diverse natural course, ALF has been sub-classified as hyperacute (IEI ≤ 7 day), acute (IEI ≤ 4 weeks) and sub-acute ALF (IEI ≥ 5 week to ≤12 weeks) by British authors. Despite these differences in definitions, the central and most important clinical event in ALF is occurrence of hepatic encephalopathy (HE) and cerebral edema which is responsible for most of the fatalities in this serious clinical syndrome. The pathogenesis of encephalopathy and cerebral edema in ALF is unique and multifactorial, and evaluation of these pathogenetic processes provides insight into its effective treatment strategy. Additionally, infection and coagulopathy have been identified to ensue rapidly in these patients - which also determines outcome with resultant management challenge.

**COMPLICATIONS AND CAUSES OF DEATH IN ACUTE LIVER FAILURE**

The major complications in ALF usually associated with death include cerebral edema, seizures, infections, bleeding due to coagulopathy and renal failure. These events infrequently get further aggravated by electrolyte and acid base imbalance and hypoglycaemia. The complications of ALF vary by region and by etiology.
Cerebral edema has been documented to be the most common cause of mortality. In India 58% of ALF patients have cerebral edema at the time of hospitalization. The mortality rate of patients with cerebral edema has been reported as 82%, compared to 44% among patients without cerebral edema. Older studies from the UK reported that overt features of cerebral edema increase in frequency with increasing grades of HE. However, with advent of intracranial pressure estimation, it is now clear that most patients with ALF at the time of hospitalization have some degree of cerebral edema and need careful monitoring. Recent data suggest that cerebral edema is less frequent now than in former years, but this may reflect earlier admission to hospital and better intensive care unit care.

Infection is a common complication in ALF that has been documented across the globe. An incidence of infection has been reported as high as 90% has been reported in the initial series from UK; the causative organisms being bacteria in 80% of cases and fungal infections in 32%. The predominant organisms are Gram-positive bacteria and the most common site of infection is respiratory tract. In more recent reports, the predominant organisms reported are Gram-negative.

Renal failure has been described in 40%–80% of patients in western series, and is associated with a poor prognosis. It occurs more frequently in acetaminophen induced ALF (70%) and less frequently in ALF due to other causes (30%) in western series, suggesting that there may be a toxic effect of acetaminophen on renal tubules. Renal failure is reported in 10% of patients from India. Gastrointestinal bleeding is reported in 7–20%. However gastrointestinal bleed has rarely been implicated as the cause of death and usually occurs as a terminal event associated with other complications.

These complications may occur at presentation, or may develop subsequently. They may occur in isolation or in combinations. Management of each complication is important as it influences the ultimate outcome. Encephalopathy and cerebral edema are common presenting features over which other complications like infections, renal failure and gastrointestinal bleed may supervene, perpetuating the brain edema and consequent death. Therefore it is important to understand the pathophysiological drivers of encephalopathy. Intervention at the level of these pathogenetic mechanisms, along with specific therapy of complications, forms the mainstay of medical management of ALF.

**PATHOGENETIC DRIVERS**

The underlying pathogenesis of encephalopathy, consequent cerebral edema and raised intracranial hypertension (ICT) in ALF is complex and has been extensively studied. Ammonia has been implicated as the major neurotoxin in ALF. In addition, Systemic Inflammatory Response Syndrome (SIRS) and loss of autoregulation of cerebral blood flow have been implicated as other important pathogenetic events in accentuating encephalopathy and cerebral edema in ALF.

**Role of Ammonia**

In animal models as well as in patients with ALF, marked swelling of astrocytes has been documented. In animal models of ALF, expression of various astrocytic proteins has been reported. Astrocytes are specialized neuroglial cells, initially considered as passive supporters of the neuronal framework, which are now recognized to play crucial roles in brain metabolism, maintaining the blood brain barrier, modulating synaptic transmission, and neural inflammation. Changes in astrocytes are documented to be due to hyperammonemia in ALF. Astrocyte swelling due to increased brain water and alteration in the functional property of the astrocyte causes encephalopathy and cerebral edema resulting in intracranial hypertension, which often results in brain herniation, the most important mechanism of death in ALF. The source of circulating ammonia is primarily derived from glutamine metabolism in the intestinal epithelium. Intestinal epithelium undergoes rapid turnover and uses glutamine as a source of energy. Intestinal epithelium contains glutaminase as well as glutamine synthase, and glutaminase converts glutamine to glutamate and ammonia. Some amount of ammonia is also generated by the urease activity of gut flora and renal production (kidney also contains glutaminase and glutamine synthase). The circulating ammonia to some extent is excreted by the kidney, used by the muscles to re-synthesise glutamine (muscle also has glutamine synthase and glutaminase), but predominantly is converted to urea (Kreb’s urea cycle present in periportal hepatocytes) as well as glutamine (by glutamine synthase present in perivenous hepatocytes) in the liver. Brain also contains glutamine synthase as well as glutaminase. Therefore, brain can synthesise glutamine from ammonia as well as metabolise glutamine to glutamate and ammonia. Thus, ammonia is predominantly metabolized in the liver and is converted to urea and glutamine. The enzymes for a complete urea cycle are exclusively localized in the liver. Urea is water soluble and is excreted in urine which is the major pathway for ammonia disposal. Thus, the ability of liver to metabolise ammonia is grossly compromised in ALF resulting in hyperammonemic state, which exerts deleterious effects contributing to HE.

The neurotoxic effect of ammonia in ALF

1. Alteration in Cell (Astrocyte) volume regulation
2. Alteration in cerebral energy homeostasis
3. Alteration in astrocytic and neuronal protein expression affecting its structure and functions
4. Oxidative and nitrosative stress
The arterial ammonia level measured within 24 h of patient presentation has also been demonstrated to correlate with patient outcome in ALF, unlike the case in cirrhosis.4,17 A study conducted by us documented that non-survivors had significantly higher median ammonia levels than survivors (174.7 v 105.0 mmol/L; \( P, 0.001 \)) in patients with ALF.4 An arterial ammonia level of \( \geq 124 \) mmol/L was found to predict mortality with 78.6% sensitivity and 76.3% specificity, and had 77.5% diagnostic accuracy. Patients with higher ammonia levels also developed more complications, including deeper encephalopathy, cerebral edema, need for ventilation, and seizures. Logistic regression analysis showed that pH, presence of cerebral edema, and arterial ammonia at admission were independent predictors of mortality (odds ratios 6.6, 12.6, and 10.9, respectively).4 However, ALF is a dynamic process where variables determining prognosis at admission change over time, and thus the clinical course varies accordingly. The ALF Early dynamic model (ALFED) is based on whether the levels of predictive variables remain persistently high or increase over 3 days above the discriminatory cut-off values. The model has four variables: arterial ammonia \( \geq 123 \) µmol/L, serum bilirubin \( \geq 15 \) mg/dl, international normalized ratio \( \geq 5 \) and hepatic encephalopathy > Grade II. These values if present on day 3 of presentation are independent predictors of mortality. The scores have been depicted in Table 1; total scores vary from 0 to 6.

The performance of the ALFED model was superior to the King’s College Hospital criteria and the Model for End stage Liver Disease score, even when their 3-day serial values were taken into consideration. An ALFED score of \( \geq 4 \) had a high positive predictive value (85%) and negative predictive value (87%) in the validation cohort.18 The most widely used prognostic model is the King’s college criteria (Table 2). However, in Indian situation, its relevance is questionable. In India paracetamol poisoning associated ALF is practically non-existent,2,4 and therefore the prognostic criteria for paracetamol poisoning is not practiced. The King’s prognostic criteria for non-paracetamol poisoning include 5 parameters (Table 2), of which 3 parameters are not prevalent in Indian situation.

<table>
<thead>
<tr>
<th>Table 1 ALFED Model.18</th>
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<tbody>
<tr>
<td>Variables over 3 days</td>
</tr>
<tr>
<td>------------------------</td>
</tr>
<tr>
<td>Hepatic Encephalopathy</td>
</tr>
<tr>
<td>Persistent or progressed to Grade &gt;2</td>
</tr>
<tr>
<td>INR Persistent or increased to ( \geq 5 )</td>
</tr>
<tr>
<td>Arterial ammonia Persistent or increased to ( \geq 123 ) mmol/L</td>
</tr>
<tr>
<td>Serum bilirubin Persistent or increased to ( \geq 15 ) mg/dl</td>
</tr>
</tbody>
</table>

Total score calculated by adding individual scores of all 4 variables.

<table>
<thead>
<tr>
<th>Table 2 ALF Prognostic Model (King’s College Criteria).</th>
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<tbody>
<tr>
<td>Non Acetaminophen ALF</td>
</tr>
<tr>
<td>------------------------</td>
</tr>
<tr>
<td>Arterial ammonia &gt;123 µmol/L or arterial pH &lt;7.3</td>
</tr>
<tr>
<td>INR &gt;6.5 (or PT &gt;100 s)</td>
</tr>
<tr>
<td>Any 3 of the following:</td>
</tr>
<tr>
<td>• Age: &lt;10 or &gt;40 years</td>
</tr>
<tr>
<td>• Etiology: Non A, Non B, drug induced</td>
</tr>
<tr>
<td>• I-E interval: &gt;7 days</td>
</tr>
<tr>
<td>• INR &gt;3.5 (or PT &gt;50 s)</td>
</tr>
<tr>
<td>• Bilirubin: &gt;17.5 mg%</td>
</tr>
</tbody>
</table>

Most Indian patients with ALF are 20–30 years old at presentation, almost all patients have an I-E interval of <7 days and Non A, Non B virus (its exact nature is not known) and drug idiosyncrasy as etiology of ALF are absent among Indian patients. Therefore the ALFED model may be more applicable in countries like India and probably in other developing countries in Asia and Africa where two third of world’s population lives.

Infections and Systemic Inflammatory Response Syndrome (SIRS) in Acute Liver Failure

Lever is an important component of the immune system. In ALF, rapid immune paralysis is well known and overt sepsis has been reported in 55%–90% of patients.17 A strong association between occurrence of infection and course of encephalopathy has been clearly documented.2,20 In a prospective evaluation of 96 acetaminophen induced ALF cases with early encephalopathy, occurrence of infection was associated with progression of encephalopathy to deeper grades in about 80% of the individuals.21 In this study, ALF patients without evidence of overt infection (\( n = 168 \)), who progressed from early to advanced encephalopathy had a number of components of SIRS.21 Fifty percent of those with 2–3 component of SIRS progressed to advanced encephalopathy in contrast to 25% of the patients without any components of SIRS who had progression of encephalopathy. The components of SIRS are highlighted in Table 3.22

Astrocytes form an integral component of the blood–brain barrier (BBB). Ammonia gets accumulated in astrocytes, theoretically altering the permeability of the BBB. It thus ‘primes’ the brain to participate in the systemic inflammation stimulated predominantly by sepsis. Endotoxins and cytokines, the major mediators of the systemic inflammatory response, are unable to cross the BBB. They
interact with receptors in the endothelial cells and astrocyte processes on the BBB and can disrupt the capillary tight junctions. They stimulate multiple downstream pathways associated with NO signalling, COX and cytokines. In the late stages of acute liver failure, pro-inflammatory cytokines are observed to efflux from the brain, indicating brain production of TNF-α, IL-6 and IL-1β during uncontrollable intracranial hypertension. This inflammatory response is considered to mediate vasodilatation and oxidative stress, both adding to the cytotoxic effects of ammonia. Even peripheral cytokines subsequent to sepsis in patients with liver failure can cross the BBB. Cytokines have been documented to increase intracranial hypertension in pig models of ALF. Liver failure increases an individual’s susceptibility to sepsis.

**Loss of Cerebral Auto Regulation**

Usually cerebral blood flow (CBF) is independent of change in systemic blood pressure and blood flow to various regions of brain varies depending upon the metabolic need of neuron (glucose and oxygen). This delicate balance of regional CBF and the metabolic requirement is lost in patient and animal models of ALF. This may cause decrease in CBF, cerebral congestion which aggravates cerebral edema. CBF correlates very closely with the brain water and ICP in experimental ammonia induced brain edema. Normally, the brain autoregulation can maintain a normal CBF at arterial pressures from 65 to 140 mm Hg. This loss of autoregulation of CBF in ALF signifies that even mildly elevated cerebral perfusion pressure (CPP) will increase CBF and hydrostatic pressure across the BBB, while episodes of arterial hypotension may stop capillary blood flow, leading to cerebral hypoxia. This phenomenon has been called ‘dissociated cerebral vasoparalysis’. CBF is variable in ALF, and may be excessive at some time point and inadequate at others. CBF is reduced in most patients with grade 3 or 4 encephalopathy, even with maintained mean arterial pressure. However, increased CBF precedes high ICP and cerebral herniation. This loss of CBF autoregulation is rare in patients with cirrhosis or sepsis. Even though the exact mechanism of loss of autoregulation of CBF is poorly understood, it is a definite event in ALF perpetuating the process of cerebral edema. This loss of autoregulation is restored within 1 day of liver transplant and within 4 days of spontaneous recovery. Even hypothermia restored cerebral autoregulation in a series of 14 patients with ALF.

**Aggravating Factors of Encephalopathy in Acute Liver Failure**

In the rat model, hyponatremia along with ammonia have been documented to induce brain edema, and hypernatremia in patients with ALF seems to protect against brain edema. About 25% of patients with ALF in a report from India had hyponatremia, which was prominent in patients with severe encephalopathy. However, pathogenesis of hyponatremia in ALF is unclear. Use of sedatives in patients with ALF presenting with agitation causing advanced encephalopathy is a common clinical experience. Alkaline pH is also known to drive ammonia in to the astrocyte in patients with hepatic encephalopathy. These factors are reversible and therefore important to be assessed in such patients.

Hyponatremia, alkalosis, SIRS have been well identified by now to increase susceptibility of astrocytes to increased arterial ammonia levels. Therefore control and prevention of these factors, along with ammonia lowering therapy, today remains the most rational approach to manage patients with hepatic encephalopathy. Figure 1 summarizes the essential mechanism of encephalopathy and cerebral edema in ALF and the possible therapeutic interventions directed towards these pathogenetic changes.

**MANAGEMENT OF ENCEPHALOPATHY IN ACUTE LIVER FAILURE**

Current specific management of encephalopathy in ALF is based on measures that reduce or eliminate ammonia, prevent sepsis and, if sepsis has set in, treat it aggressively. While these rationales remains the mainstay of therapeutic approach, treatment in an intensive care unit with availability of organ support systems to prevent death due to organ failure also continues to be the other important aspect in the management of hepatic encephalopathy in ALF. The essential steps of management of hepatic encephalopathy are depicted in Table 4.

However certain details of specific support in encephalopathy in ALF need to be discussed in greater detail and can be categorized as 1) medical therapy directed to control encephalopathy and cerebral edema 2) ammonia lowering strategies 3) other forms of therapy directed to control and prevent complications like hemodynamic instability, renal failure, coagulopathy, hypoglycaemia and respiratory arrest. 4) liver transplant and bioartificial liver support systems.

**Medical Therapy Directed to Control Encephalopathy and Cerebral Edema**

**Neurologic Support**

All patients should be nursed in a 20°–30° head up tilt to improve venous drainage. Patient turning and other tactile stimulation should be minimized. Respiratory suctioning should be limited to less than 15 s duration at a time. One to two ml of lidocaine instilled in the endotracheal tube and ventilation with 100% oxygen prior to suctioning may be helpful in preventing surges of raised ICP. Control of agitation is important, as valsalva maneuver from psychomotor agitation may lead to elevation of the ICP. A diligent
search for treatable causes of agitation like distended bladder, and bedsores should be made (Table 5). Small intravenous doses of short acting benzodiazepines should be reserved as a last resort for serious psychomotor agitation.

**Intracranial Pressure Monitoring**

Traditional signs of elevated ICP are unreliable in ALF. Computed tomography of the head is not a reliable way to estimate ICP in ALF. Currently many centres install an extradural ICP monitoring device once the patient has developed grade 3 or 4 encephalopathy. Prior endotracheal intubation is necessary under propofol anesthesia. Aggressive correction of coagulopathy and thrombocytopenia must be undertaken with a goal of achieving a platelet count of >50,000/mm$^3$ and an INR <1.7 or PT within 2–3 s of normal. Correction of the PT usually requires bolus administration of 2 units of fresh frozen plasma (FFP) followed by a continuous FFP infusion at a rate of 75–150 ml/h. The transducer to measure the ICP is placed by a neurosurgeon in the ICU under local anesthesia. The standard site is the right frontal area chosen to avoid the (right handed) dominant lobe, sagittal sinus, and motor cortex.

Epidural transducers are used most commonly and carry the lowest complication rate (3.8%). Subdural bolts and parenchymal monitors carry complication rates (mainly bleeding) of 20% and 22% respectively. Colonization of ICP devices increases significantly after about 5 days.

![Figure 1 Mechanism of encephalopathy in ALF and therapeutic intervention directed to these pathogentic mechanisms.](image)

**Table 4 Principles in the Management of Hepatic Encephalopathy.**

1. **Aggressive supportive therapy:** Intensive care unit, Organ support system, Nutrition
2. **Identification and removal of precipitating factors:** Control of GI bleed, Sepsis, Hyponatremia, renal failure, Constipation, Psycho-active drugs.
3. **Reduction of Nitrogenous load from the Gut:** Lactulose, Antibiotics, Enema
4. **Manipulation of neurotransmitters:** Flumazenil, Branched chain amino acids
5. **Ammonia Lowering Therapy:** L-Ornithine L-Aspartate (LOLA), L-Ornithine Phenyl Acetate (LOPA), Sodium Benzoate, Hypothermia.
6. **Novel therapies:** Molecular Adsorption Recirculating System (MARS), Probiotics with increased capacity to consume Ammonia

**Table 5 Factors that Increase Intracranial Pressure.**

<table>
<thead>
<tr>
<th>Factor</th>
<th>Cause</th>
</tr>
</thead>
<tbody>
<tr>
<td>Valsalva maneuver</td>
<td>Arterial hypertension</td>
</tr>
<tr>
<td>Positive end expiratory pressure (PEEP)</td>
<td>Vomiting, shivering</td>
</tr>
<tr>
<td>Head and body turning or moving</td>
<td>Psychomotor agitation</td>
</tr>
<tr>
<td>Neck veins compression</td>
<td>Noxious stimuli</td>
</tr>
<tr>
<td>Respiratory suctioning</td>
<td>Seizures</td>
</tr>
<tr>
<td>Fever</td>
<td>Vasodilatory agents</td>
</tr>
<tr>
<td>Horizontal decubitus</td>
<td>Isometric muscle exercise</td>
</tr>
<tr>
<td>Severe hypoxemia</td>
<td>Trendelenburg position</td>
</tr>
<tr>
<td>Any degree of hypercapnia</td>
<td>Coughing, sneezing</td>
</tr>
</tbody>
</table>
of insertion, and the risk of superficial wound infection and meningitis rises.

In adults, the average ICP ranges from 0 to 10 mmHg. The maximal permissible upper limit is 20 mmHg. The overall goal is to maintain an ICP of less than 20 mmHg and CPP (MAP–ICP) above 50 mmHg at all times. A slow progressive increase in ICP to >25 mmHg, or pressure waves rising to 30–50 mmHg lasting for 5–20 min, reflect falling cerebral compliance. Prolonged (>2 h) elevation of ICP to >40 mmHg or a reduction in CPP < 50 mmHg are considered a contraindication for liver transplantation.

**Reverse Jugular Venous Monitoring**

Jugular bulb venous saturation is frequently used in neuro-surgical intensive care, and involves insertion of a fine catheter in a retrograde fashion until its tip lies in the jugular bulb. Blood returning from the cerebral circulation can then be sampled. A jugular venous saturation of <55% represents an ischemic brain, while a saturation of >85% represents a hyperemic brain.37 Measures to improve oxygen supply are indicated in the former situation, and measures to reduce cerebral blood flow in the latter situation.

**Osmotherapy**

Mannitol lowers the ICP by reducing the brain water and changing the rheological characteristics of blood. When the ICP rises to over 20–25 mmHg for more than 5 min, an intravenous bolus of mannitol should be given (0.5–1 g/kg, 20% solution, over 5 min). Repeated boluses may be administered as long as the serum osmolality is <320 mOsm/L. The response to a mannitol bolus may be expected 15–60 min post-injection. In about 20% of patients a paradoxical increase in ICP occurs after mannitol infusion.38 High doses can result in acute renal failure and damage to the BBB. Mannitol works best in mild to moderate intracranial hypertension and is less effective when the ICP is greater than 60 mmHg.39 In oliguric/anuric patients, mannitol should only be given in combination with continuous veno venous or arteriovenous hemodiafiltration.

**Thiopentone**

Based on data suggesting that barbiturates could be of value in controlling the intracranial hypertension of head injury, intravenous thiopental was assessed in 13 patients with fulminant hepatic failure complicated by unresponsive intracranial hypertension. The ICP was reduced in all cases, and in eight cases thiopentone infusion achieved stable normal intracranial and cerebral perfusion pressure. Five patients made a complete recovery.40 The recommended dose of pentobarbital is a loading dose of 3–5 mg/kg (maximum 500 mg) over 15 min, followed by a continuous infusion of 0.5–2.0 mg/h. Barbiturate therapy must be used with simultaneous continuous ICP and arterial blood pressure monitoring.

**Hypothermia**

Moderate hypothermia (32 °C–35 °C) using cooling blankets leads to a reduction in CBF, cerebral metabolism, ammonia uptake by the brain, and glutamine synthesis, and reduces intracranial pressure in patients with ALF.41 Data from studies in patients undergoing liver transplantation for ALF suggests that an increase in intracranial pressure can be prevented during the dissection and reperfusion phases of the operation if the patients are maintained hypothermic during surgery.42 However hypothermia as a therapy has not shown any improvement in survival among patients with ALF but has been used as a therapeutic bridge to transplantation.43 Hypothermia in the above mentioned studies has been used for not more than 8 h, and prolonged hypothermia is well known to cause cardiac arrhythmia and coagulopathy. Therefore routine use of hypothermia for short duration may be of help particularly in patients with ALF.

**Prophylactic Phenytoin**

It has been suggested that subclinical seizure activity in patients with deep encephalopathy on ventilation may remain unrecognized and may lead to exacerbation of cerebral hypoxia and edema. Hence it was recommended that all patients with grade III or IV encephalopathy should be monitored for subclinical seizure activity and treated with prophylactic phenytoin. However, in a randomized, controlled clinical trial of 42 patients, it was found that prophylactic use of phenytoin did not prevent cerebral edema, seizures or improve survival.44

**Hyperventilation**

Patients with ALF often hyperventilate spontaneously, and attempts to control this are of no use. Hyperventilation reduces CBF, and may be useful in the subgroup of patients with cerebral hyperemia as reflected by jugular bulb venous saturation of more than 75%. Prophylactic hyperventilation does not reduce the frequency of intracranial hypertension in ALF, but a moderate reduction in pCO2 to 25–30 mmHg is helpful in decreasing the ICP once cerebral edema has begun to develop.45 Excessive hyperventilation may lead to cerebral vasoconstriction.

**Ammonia Lowering Strategies**

Benefits of lactulose have been seriously questioned. In a recent meta-analysis, only few high quality studies have shown any beneficial effect with lactulose and its popularity stems from a long clinical experience rather than strong objective evidence.46 Antibiotics have their inherent adverse effects, especially on prolonged use. Concepts on protein restriction are being revised.47 Newer agents to reduce ammonia like phenyl acetate and probiotics, having capacity to consume ammonia, are being introduced.
Novel approaches like L-carnitine, flumazenil in deep coma even when not precipitated by benzodiazepines is being vigorously investigated.

**L-Ornithine L-Aspartate (LOLA):** LOLA is a stable salt of the amino acids ornithine and aspartic acid. These two amino acids get converted to glutamate in the muscles and hepatocytes. Glutamate is the substrate on which the enzyme glutamine synthase (present in the muscle as well as liver), acts and combines it with ammonia to make glutamine and thereby reduces blood ammonia levels. In liver failure, the enzyme glutamine synthase in the muscle is up-regulated, and therefore it is hypothesized that LOLA may reduce arterial ammonia levels. It has been shown that LOLA reduces arterial ammonia levels accompanied by improvement in mental status in patients with hepatic encephalopathy associated with chronic liver disease. In rat model of acute liver failure (ALF), LOLA has been shown to prevent brain edema and hepatic encephalopathy which was accompanied by up regulation of muscle glutamine synthase activity and plasma glutamine levels were also raised indicating that LOLA in ALF may be beneficial. A randomized placebo controlled double blind trial of LOLA in patients with ALF was recently published. However LOLA therapy was not associated with decrease in ammonia levels or improvement in encephalopathy or survival (Figure 2).

In the above mentioned study ammonia levels were measured for 6 days among patients receiving LOLA as well as placebo, however the rate of decline of ammonia was also similar between the two arms (Figure 3). Peculiarly, patients receiving LOLA had more frequent seizures. Very high glutamine levels in the systemic circulation are found in ALF. LOLA could theoretically further increase ammonia detoxification by the skeletal muscle by increased glutamine synthesis. This glutamine is recycled back to the intestine and kidney where it is broken down to ammonia and glutamate by glutaminase, thus LOLA is ineffective in reducing the ammonia levels.

**L-Ornithine Phenyl Acetate (LOPA):** Brusilow in 1979 suggested use of alternative pathway for ammonia disposal in Urea Cycle Disorders (UCD). The drugs proposed were phenyl acetate and sodium benzoate. Phenyl acetate combines with glutamine to form phenylacetylglutamine which is water soluble and is excreted in urine. One molecule of phenyl acetate removes two molecule of Glutamine and this became the mainstay of management in UCD in the ensuing time.

Recently Davies et al from the university college of London (UCL) documented that in bile duct ligated rat model of cirrhosis, LOPA could reduce arterial ammonia, associated with decrease in cerebral glutamine/myoinositol ratio, brain water content and increase in arterial glutamine concentration along with increase in urinary phenylacetylglutamine levels. Ytrebo et al from the same UCL group in a devascularised pig model of ALF documented that LOPA prevented rise of plasma and brain ammonia concentration and prevented rise in intracranial pressure. This latter study also documented increase in urinary phenylacetylglutamine excretion along with increase in muscle glutamine synthetase activity. Ornithine has been used in these two studies to induce glutamate formation in the muscle. However whether these agents will be effective in humans needs evaluation. Further, a phenyl acetate blood...
level exceeding >6 mmol/L is associated with toxicity and in presence of renal failure its use will be limited.

**Sodium benzoate:** Sodium Benzoate is usually used as a food preservative. It combines with glycine to form hippurate which is water soluble and gets excreted in urine easily and thereby may decrease arterial ammonia levels. However, therapeutic trials in humans show conflicting results. It has never been used in patients with acute liver failure. However our own experience does not support its ammonia lowering effect in patients with ALF.

**Rifaximin:** Rifaximin is an oral antibiotic with a broad spectrum of activity against enteric bacteria. It is minimally absorbed from the gut (<0.4%). Safety profile and tolerability is comparable to placebo. No dose adjustment is required in patients with renal insufficiency. Current data suggests that rifaximin is as effective as established therapies like lactulose or neomycin in treatment of HE and has a better safety profile and tolerability in patients with chronic liver disease. Its’ possible benefit in lowering ammonia in patients with ALF has not yet been explored.

**Probiotics:** Probiotics promote growth of non-urease producing bacteria in the colon thereby reducing generation and absorption of ammonia by suitably altering the gut flora. Probiotics have been shown to reduce ammonia and improve cognitive function in patients with minimal HE. There is no data on its role in ALF.

**Miscellaneous interventions to lower ammonia and improve encephalopathy:** In a recently published trial by USALF study group, 21 days transplant free survival was documented to be significantly better in the NAC treated than the placebo treated patients, however overall survival between the treated and control group was similar. In a controlled trial of 44 patients it was found that the use of dexamethasone could not prevent the development of cerebral edema or improve survival. Any unexplained decrease in blood pressure, drop in urinary output, worsening encephalopathy, and development of severe acidosis or disseminated intravascular coagulation should be considered as signs of sepsis. Fungal infection should be strongly considered in the setting of unresponsive fever, leukocytosis, and deterioration of neurologic status after initial improvement and presence of renal failure. GM-CSF has been reported to improves neutrophil function both in vitro and in vivo among ALF patients.

**Hemodynamic Support**

ALF patients have a high cardiac output, low systemic vascular resistance, and relative arterial hypotension. An adequate cardiovascular filling pressure (PCWP: 8–14 mmHg) must be assured. The goal is to maintain the mean arterial pressure (MAP) above 60 mmHg. Vasoressor agents are indicated if the MAP is <60 mmHg despite adequate intravascular volume. Oxygen consumption may decrease with the use of vasoressor agents despite the increased arterial pressure, as a result of reduction in oxygen delivery and extraction rates. Reduced oxygen extraction may be prevented with concurrent use of prostacyclin. Arterial hypotension in spite of an adequate intravascular volume should be considered a sign of bacterial or fungal sepsis and treated accordingly.

**Coagulopathy**

The coagulopathy of ALF may be clinically silent or manifested by bleeding from mucosal membranes. Infusion of FFP is indicated only for active bleeding or before invasive procedures. The risk of bleeding from stress ulceration of gastric mucosa is reduced by the prophylactic use of sucralfate.
Acute Liver Failure

Mechanical Ventilation of Patients with Liver Failure

The lungs are relatively spared in ALF and standard modes of mechanical ventilation can be used. Most patients will tolerate ventilation with minimal sedation. The risks of paralysis may outweigh the benefits in the majority of patients, but may have a role in selected patients with refractory intracranial hypertension. Barbiturates and midazolam produce a dose-dependent reduction in metabolic rate, CBF and cerebral blood volume. A traumatic intubation may induce a sudden rise in the ICP and provoke herniation. Hence tracheal intubation must be gentle. Standard ventilation modes, most commonly synchronized intermittent mandatory ventilation (SIMV) are used. The usual initial ventilator settings are: respiratory rate of 14–18 breaths/minute; tidal volume of 10–12 times the body weight in kilogram; inspiratory period of 25%–33%; pause time of 10% of total cycle; and PEEP of 2–4 mmHg. High levels of PEEP (more than 10–15 mmHg), lead to worsening or facilitating the development of cerebral edema, and a decrease in hepatic arterial blood flow. Propofol inhibits sympathetic vasoconstrictor activity and has negative inotropic effect. Among the muscle relaxants, pancuronium, vecuronium, and rocuronium exhibit a prolonged elimination in the presence of liver failure. Atracurium is degraded by the non-enzymatic Hoffman elimination, which may reduce CPP by increasing cerebral edema through rapid osmolar shifts or reducing the MAP. Dialysis reduces ammonia levels along with blood urea and glutamine.

Nutritional and Metabolic Support

Almost half of ALF patients develop hypoglycaemia. Development of hypoglycaemia can be sudden and may confound the interpretation of mental changes. Glucose requirements in these patients are highly variable and require close monitoring. Blood sugar should be monitored at 2–3 h intervals. Whenever the blood glucose level is lower than 60 mg/dl, an intravenous bolus of 50–100 ml of 50% dextrose should be administered. The amount of water administered as a solvent for dextrose should be minimized by providing solutions concentrated to 25%–50%.

Recent evidence suggests that the glucose transport across the blood brain barrier is increased because of upregulation of glucose carriers in ALF patients. Hyperglycaemia may contribute to raised ICP because increased glucose influx leads to cerebral lactic acid accumulation. Hence it may be prudent to keep blood glucose within the normal limits.

ALF is a hypercatabolic state. Energy requirements in ALF are increased by as much as 60% and are further elevated by the presence of complicating infection. Whole body protein catabolism may be increased up to four times the normal rate. Massive amino acid losses in the urine occur. A combination of parenteral dextrose and lipid emulsions, and at least 40 gm protein/day should be administered initially. It has been shown that lipid emulsions may be used safely in patients with ALF. There is no reason to restrict proteins in ALF. Branched chain amino acids (BCAA) offer no additional advantage, with the exception of the patients requiring frequent dialysis, in whom large BCAA losses may occur. Hypokalemia, hypomagnesemia, hypophosphatemia and hypocalemia are common and must be corrected. Either the enteral or parenteral route may be used for feeding, but the former route has obvious advantages.

Liver Transplantation and Bio-artificial Liver Support

Liver Transplantation

Overall, the 1 year survival after cadaveric orthotopic liver transplantation (OLT) ranges from 50% to 75%. In experienced centres, outcome with split liver grafts is comparable with that after use of full size organs. Survival is substantially reduced among patients with sepsis and multiorgan failure before OLT. Auxiliary partial OLT, as a form of temporary liver support in patients with ALF, has been reported to have results similar to those of conventional transplantation. Withdrawal of immunosuppression, leading to graft atrophy may be possible in up to 65% patients surviving 1 year. Living donor liver transplantation has
been more often used in children. The 1 year survival rate after living related OLT in a total of 35 paediatric cases in 3 series was 59%–90%.73–75 The 1 year survival in the largest series of living related OLT in adults, including 53 patients has reported a 75% survival at 1 year.76

**Non Transplant Therapies for Liver Support**

Therapeutic interventions that have been used to treat ALF in lieu of OLT are listed in Table 6.

Extracorporeal Liver Support may be broadly classified as artificial which contain no biologic component (e.g. MARS), or bioartificial which include viable liver cells in culture within bioreactors or involve perfusion of the patients’ blood through an isolated human, or porcine whole liver. However, none of these bioartificial support system till date has been shown to improve survival among patients with ALF.

**Molecular Adsorbent Recirculating System (MARS)**

MARS (Teraklin AG, Rostock, Germany) is a dialysis treatment which uses a recirculating dialysate containing 20% albumin that is regenerated on line by dialysis against a bicarbonate buffered dialysate, followed by passage through a column with uncoated charcoal and a second column with an anion exchange resin. This allows the removal of both water soluble substances, such as urea, creatinine, and ammonia, and of albumin bound substances, such as phenol, bile acids, bilirubin, BCAA, and short chain fatty acids. MARS treatment can also remove cytokines like TNF-α and IL-6. MARS therapy has been commercially available since 1999, and 38 patients with ALF have been reported to the International MARS registry till 2002.77 Majority of these cases were drug induced. Overall 19 patients survived, of whom 6 were transplanted. Preliminary data shows that MARS treatment improves encephalopathy, and leads to a reduction in bilirubin and ammonia levels.78 However no randomized trials of MARS therapy in ALF are available. At our centre we conducted a randomised study comparing MARS with conventional management in patients with ALF (n = 20) and could not document any survival benefit of MARS over the conventional management (unpublished observation).

**Treatment of underlying etiology:** The management of ALF is supportive. In some specific etiologies, drugs have been shown to be effective. In patients with HBV-ALF, antivirals have been recommended. The choice of antivirals is between tenofovir and entecavir, the former being preferred in patients with renal failure due to its safety profile. The doses of antivirals need to be modified as per the creatinine clearance. In patients with antituberculosis drug induced or any other suspected drug induced ALF, the suspected drug should be immediately stopped. In HEV-ALF, presently there is no data on the use of Ribavirin, though this has been shown to be effective in managing chronic HEV in solid organ transplant patients and selected patients with HEV related ACLF.79,80 In patients with paracetamol toxicity, activated charcoal has been shown to be effective in gastric decontamination. High doses of NAC are recommended in such patients and should be started within 48 h of ingestion.81

**Pregnancy and acute liver failure:** There is increased risk of maternal and fetal complications in pregnant patients with ALF.82 The principles of management of abortions, preterm labor, premature rupture of membranes and still birth remain unchanged. Termination of pregnancy is not indicated. There is an increased risk of bleeding due to associated coagulopathy. Management of pregnant patients with ALF is similar to other ALF patients. Complications need to be managed on a case to case basis. A detailed management of ALF in pregnancy has been published in a previous issue of this journal.81

### CONCLUSION

Hepatic encephalopathy in ALF is its protean manifestation. Hyperammonemia, SIRS, loss of cerebral autoregulation, hyponatremia and imbalance in acid-base homeostasis are major pathogenetic mechanism of encephalopathy in ALF. Cerebral edema and infection are the major causes of death in this disease. Specific strategy for neurological support, prevention of infection and its treatment, ammonia lowering therapies, maintenance of acid-base balance, close monitoring in an ICU setup with adequate support for failing organs remains the mainstay of medical management of encephalopathy in ALF. Patients with ominous prognostic criteria benefit from liver transplant. However liver support devices have not been very beneficial in ALF till date.

### CONFLICTS OF INTEREST

All authors have none to declare.
REFERENCES


