Acute-on-chronic liver failure in liver transplant candidates with non-alcoholic steatohepatitis

Iliana Doycheva, Paul J. Thuluvath

Institute for Digestive Heath and Liver Disease, Mercy Medical Center, Baltimore, MD, USA

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Correspondence to: Paul J. Thuluvath, MD. Institute for Digestive Health and Liver Disease, Mercy Medical Center, 301 St. Paul Place, Baltimore, MD 21202, USA. Email: thuluvath@gmail.com.

Abstract: Non-alcoholic fatty liver disease (NAFLD) has become the most common liver disease worldwide. It is expected that non-alcoholic steatohepatitis (NASH), NASH-related cirrhosis and its decompensated forms will increase further in the next two decades. Acute-on-chronic liver failure (ACLF) is a distinct syndrome characterized by rapid deterioration of liver function in patients with chronic liver disease that is associated with development of one or more organ failures, and carries a very high short-term mortality. There is a paucity of data on ACLF in patients with NASH cirrhosis. Recent studies have shown that although ACLF incidence due to NASH is lower when compared to other etiologies, NASH is the fastest growing liver disease etiology among all ACLF hospitalizations. Higher rates of infections, as a precipitating factor, and circulatory failure were noted in this population. Metabolic derangements such as obesity and diabetes might also play a confounding role in the pathophysiology, clinical course, and prognosis of NASH patients with ACLF. Patients with ACLF due to NASH have shown a lower inpatient mortality despite a longer hospital length-of-stay and a higher 28- and 90-day mortality. Patients with ACLF should be promptly transferred to a transplant center and evaluated for liver transplantation (LT). Optimal prognostic scores, timing of LT, and the best bridge to LT therapy and treatment of post-LT complications need to be elucidated in prospective studies.

Keywords: Acute-on-chronic liver failure (ACLF); non-alcoholic fatty liver disease (NAFLD); non-alcoholic steatohepatitis (NASH); survival

Introduction

Non-alcoholic fatty liver disease (NAFLD) and its aggressive form nonalcoholic steatohepatitis (NASH) has become the most common liver disease worldwide with an estimated global prevalence of 25% (1). The disease burden is attributed to obesity and metabolic syndrome and mirrors the rapidly rising global prevalence of obesity in both children and adults (2) as well as the upward trend of type 2 diabetes that occurs despite implementation of preventive measures (3). It has been estimated that approximately two-thirds of adults with obesity and 58% of diabetics worldwide have NAFLD (1). The NAFLD is forecasted to increase to 33.5% in 2030 in the USA with proportionally growing incidence of decompensated cirrhosis and hepatocellular carcinoma due to NASH (4). This will result in more patients with decompensated liver disease that will require hospitalizations and will eventually need liver transplantation (LT).

A subpopulation of patients with chronic liver disease and/or decompensated cirrhosis may develop acute-on-chronic liver failure (ACLF), a distinct syndrome characterized by rapid deterioration of liver function,
development of one or more organ failures (OFs), and high short-term mortality (5). There is significant variability in definitions of ACLF with more than ten existing ones that have used different clinical, laboratory, and prognostic criteria, which largely restricts comparability between studies (6). Currently, there are three major definitions including the one of the European Association for the Study of the Liver-Chronic Liver Failure (EASL-CLIF) that is based on the results of a large, multicenter, prospective, observational study (CANONIC) (7); the recently updated consensus definition of ACLF of the Asian Pacific Association for the Study of the Liver (APASL) (8); and the definition of the North American consortium for the study of end-stage liver disease (NACSELD) (9) (Table 1).

<table>
<thead>
<tr>
<th>Organization</th>
<th>Derivation cohort</th>
<th>Definition</th>
<th>Score and its components</th>
<th>Definition of organ failure</th>
</tr>
</thead>
<tbody>
<tr>
<td>EASL-CLIF (7)</td>
<td>CANONIC study prospective, 1,343 patients</td>
<td>Acute decompensation of cirrhosis (ascites, HE, GI bleed and/or infection) associated with OF and high 28-day mortality (&gt;15%)</td>
<td>CLIF-C OF: bilirubin, INR, creatinine, HE, circulatory, and respiratory system; range, 6–18</td>
<td>Bilirubin &gt;12 mg/dL; creatinine &gt;2 mg/dL; HE grade 3–4; INR &gt;2.5; use of vasopressors; PaO₂/FiO₂ &lt;200</td>
</tr>
<tr>
<td>APASL (8)</td>
<td>Consensus and observational, prospectively, enrolled 5,228 patients</td>
<td>Acute hepatic insult manifesting as jaundice and coagulopathy, complicated within 4 weeks by ascites and/or HE in previously diagnosed or undiagnosed CLD and high 28-day mortality</td>
<td>AARC: total bilirubin, INR, creatinine, HE, plasma lactate, range 5–15</td>
<td>Bilirubin ≥5 mg/dL; INR ≥1.5</td>
</tr>
<tr>
<td>NACSELD (9)</td>
<td>Prospective study, 507 patients</td>
<td>Two or more OFs due to acute infection in patients with cirrhosis</td>
<td>NACSELD-ACLF: at least 2 out of 4 OFs: HE, renal failure, respiratory failure, shock</td>
<td>HE grade 3–4; need for RRT, need for BiPAP or mechanical ventilation, need for pressors, MAP &lt;60 mmHg or SBP &lt;40 mmHg despite fluids</td>
</tr>
</tbody>
</table>

EASL-CLIF, the European Association for the Study of the Liver-Chronic Liver Failure; HE, hepatic encephalopathy; GI, gastrointestinal; OF, organ failure; ACLF, acute-on-chronic liver failure; WBC, white blood cell; PaO₂/FiO₂, ratio of arterial oxygen pressure to fractional inspired oxygen; APASL, the Asian Pacific Association for the Study of the Liver; AARC, APASL-ACLF research consortium; BiPAP, bilevel positive airway pressure; CLD, chronic liver disease; INR, international normalized ratio; NACSELD; the North American consortium for the study of end-stage liver disease; MAP, mean arterial pressure; SBP, systolic blood pressure.

Precipitating factors and pathophysiology of ACLF

Triggers of decompensation

A major difference exists between the three main definitions regarding precipitating events. The most common precipitating events for EASL-CLIF and NACSELD are bacterial infections (7,9), alcohol use, and gastrointestinal bleeding (7), while the major triggering event for ACLF in APASL consensus is hepatic insult, such as hepatitis B reactivation, drug-induced liver injury, superimposed hepatitis A or E, or alcohol consumption (10). It is important to note that 43.6% of the patients in the CANONIC study did not have an identifiable precipitating event (7).

General concepts of ACLF pathophysiology

Current data suggest that the most important underlying pathogenic mechanism of ACLF is systemic inflammation (11). This hypothesis was initially supported by the results of the CANONIC study that showed patients with ACLF had elevated leucocyte count and C-reactive protein even if they did not have underlying infection or prior history of acute decompensation (7). When patients with decompensated cirrhosis were compared to healthy controls, systemic inflammation was more severe in those with ACLF compared to those without and its severity correlated with ACLF grade (11). Chronic inflammation
in decompensated cirrhosis is attributed to translocation of pathogen-associated molecular patterns (PAMPs) that are recognized by different pattern recognition receptors: the membrane-bound toll-like receptors and the cytosolic nucleotide-binding oligomerization domain-like receptors, which leads to activation of inflammatory cascades and secretion of proinflammatory cytokines and reactive oxygen species (12). Systemic inflammation can exacerbate the preexistent systemic circulatory dysfunction and this may cause hypoperfusion and OFs leading to hepatic and extrahepatic cell damage and necrosis with resultant release of damage-associated molecular patterns (DAMPs), which stimulate the innate immune response and perpetuate acute inflammation (13). Liver injury further exacerbates systemic inflammation by cytokine/chemokine spillover to the blood and release of DAMPs.

Another important aspect of the pathogenesis of ACLF is cirrhosis-associated immune dysfunction. It results in altered homeostasis between systemic inflammation and hepatic immune defense. The latter is attributed to loss of surveillance and decreased liver synthetic function with lower production of albumin, complement, and acute phase proteins (14). While the proinflammatory phenotype is considered characteristic for the early phases of cirrhosis, the immunodeficient state is predominant in the later decompensated cirrhosis stages and ACLF. Moreover, genetic factors might be also playing a role as recent study identified two protective polymorphisms of interleukin-1 gene cluster that were associated with lower short-term mortality and lower susceptibility to ACLF (15).

**Suggested ACLF pathophysiology in patients with NASH**

The exact pathogenic factors contributing for progression of decompensated NASH cirrhosis to ACLF remain unclear. Given the majority of NASH patients also have obesity and type 2 diabetes mellitus (T2DM), these metabolic derangements are intricately interrelated and are likely to contribute for this progression. There is growing evidence on the pivotal role of gut-liver axis for the maintenance of metabolic homeostasis. It has been demonstrated that patients with obesity and NASH have a higher prevalence of small intestinal bacterial overgrowth (16). NASH has been associated with increased intestinal permeability (17). Patients with obesity and T2DM were noted to have significant gut dysbiosis with higher level of opportunistic pathogens such as Staphylococcus aureus, Escherichia coli, and Clostridium genus, while beneficial short chain fatty acid-producing bacteria were reduced (18). Moreover, animal data have shown that obesity and diabetes are related not only with translocation and elevated plasma levels of bacterial fragments such as lipopolysaccharides, but also of live commensal intestinal bacteria to blood and adipose tissue that causes persistent low-grade inflammation (19). All these factors might be contributing for development of ACLF in NASH patients.

**ACLF in NASH patients**

**Prevalence and incidence**

The prevalence of ACLF among patients with decompensated cirrhosis is estimated to range between 24% and 34% (7,9,20,21). In the CANONIC study, 10.8% of the hospitalized patients developed ACLF within 28 day (7) while 14% of stable outpatients with cirrhosis were diagnosed with ACLF within 1 year (22). These studies used EASL-CLIF diagnostic criteria to define ACLF. However, a recent retrospective study by Mahmud et al. noted significant disagreement between EASL and APASL criteria when both definitions were applied on the same population of 80,383 patients of the Veteran Health Administration (VA) system with compensated cirrhosis (23). The incidence of ACLF among NASH cirrhosis patients was estimated to be 3.4/1,000 (95% CI, 2.9–4.0) based on EASL-CLIF criteria and 18.6/1,000 (95% CI, 17.4–19.9) based on APASL criteria. When stratified by etiology of liver disease, patients with NASH and those with hepatitis C had the lowest incidence rate, but the highest 28- and 90-day mortality (51% and 73.5% by APASL and 45.2% and 59.2% by EASL definition, respectively). Infection was the most common precipitant factor for NASH patients with ACLF based on EASL definition. Among those with ACLF grade 3 (ACLF-3), patients with NASH and hepatitis C etiology had the highest rates of cirulatory failure although kidney failure remained the leading OF.

**Hospitalization rates and clinical characteristics of ACLF in patients with NASH**

A recent population-based analysis that used the National Inpatient Sample (NIS) between 2006 and 2014 noted that NASH was the fastest growing liver disease etiology of all ACLF hospitalizations with an increase of 63% over the study period (3.5% in 2006–2008 to 5.7% in 2012–2014, P<0.001) (24). There was no change in ACLF hospitalizations due to viral liver diseases, but the increase

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Natural history, prognosis, and mortality

Predictors of NASH progression to ACLF

A long-term follow up study has shown that 25% of patients with NASH progressed to cirrhosis over nine years with 10% developing decompensating events over 13 years (27). Obesity and T2DM, frequently associated with NAFLD, also contribute for liver disease progression. Clinical studies have demonstrated that obesity, an independent predictor of cirrhosis decompensation (28), is associated with infections in decompensated cirrhosis (29), and a higher in-hospital mortality in cirrhotic patients with sepsis (30). Importantly, morbid obesity [hazard ratio (HR) 1.21, 95% CI, 1.07–1.37] and diabetes (HR 1.15; 95% CI, 1.06–1.23) were linked to increased risk of dropout of NASH patients on the LT list (31).

Prognostic scores and predictors of severity of ACLF in patients with all etiologies

Different scores and prognostic markers have been used in various studies that included patients with ACLF (Table 1). Based on a derivation cohort of the CANONIC study, a CLIF-C acute decompensation score has been created that predicts 3- and 12-month mortality in hospitalized patients with cirrhosis, but without ACLF (32). This score performs better than Child-Pugh, Model for End-Stage Liver Disease (MELD), and MELD-Sodium (MELD-Na) scores. It includes age, serum sodium, white blood cell (WBC) count, serum creatinine, and international normalized ratio (INR) and ranges between 0 and 100 points with score 60 identifying a high-risk group of patients with 90-day mortality exceeding 30% and close to the 90-day mortality of ACLF-1 group in the CANONIC study (7). Among prospectively followed ACLF patients, ACLF grade at days 3–7 predicted mortality better than ACLF grade at diagnosis (33). The best independent predictors of severe early course were CLIF-C ACLF score (odds ratio [OR] 1.11; 95% CI, 1.07–1.15) and presence of liver failure defined as total bilirubin 12 mg/dL (OR 2.82; 95% CI, 1.72–4.63). This study also reported that patients with ACLF-3 at day 3–7 who had 4 or more OFs and CLIF-C ACLF score >64 carried a very poor prognosis with 100% mortality at three months. When compared to other scores, CLIF-C ACLF also best predicted 28-day mortality with a threshold of 370 points at 48 hours portending 100% mortality (34). Future studies should validate the applicability and utility of these scores in patients with NASH cirrhosis and ACLF.

Short-term mortality in NASH patients with ACLF

The estimated short-term mortality was higher in NASH patients who develop ACLF when compared to that of the overall population in the CANONIC study (7) and a
recent study that included large cohort of VA patients (21) (Table 2). However, this comparison is approximate as these populations might have consisted of patients with different proportions of ACLF severity based on ACLF grade and type of OF, age, gender, ethnicity, and comorbidities.

Role of LT

Short-term mortality of the waitlisted patients with ACLF and all etiologies

Growing evidence suggests that patients with ACLF-3 should be considered early for LT (35) and perhaps prioritized (36). If properly selected and transplanted early, these patients have post-LT outcomes comparable to the general population (37). A recent analysis of the United Network for Organ Sharing (UNOS) database demonstrated that patients with ACLF-3 and MELD-Na <25 were more likely to die or to be removed from the LT list when compared to the rest of ACLF patients (35). Another analysis of the same data registry showed that the probability of survival for more than 30 days on the transplant list is inversely related to the number of OFs with 80% removed with two OFs and 92–98% removed with three or more OFs, the latest having median time to death of 6–10 days (37). Moreover, a study that compared short-term waitlist mortality and delisting between patients with acute liver failure who were listed as status 1a and registrants with ACLF-3 at listing noted that patients with ACLF-3 had higher risk of 14-day mortality across all MELD-Na subgroups (36). Both groups in this study had numerically similar 1-year post-LT survival. The authors suggested that alternative scores for evaluation might be needed in these patients as MELD-Na does not reflect OFs.

Post-LT outcomes and predictors of survival of patients with ACLF and all etiologies

Previous studies have shown one-year post-LT survival in patient with ACLF ranging between 46% and 87% (33,38,39). However, these results are based on very small sample sizes, different ACLF definitions, and varying severity of ACLF at inclusion and at the time of transplant. One retrospective and two UNOS-based studies assessed outcomes in patients with ACLF-3 (35,37,40). One-year mortality in this population was 81-84%. Although respiratory failure requiring mechanical ventilation at the time of LT was identified as an independent risk factor in one of those studies (35), there was no association in another study (37). Other independent negative predictors for survival were high-risk donors (35,37), hepatitis C infection etiology or hepatocellular carcinoma, and the number of OFs (37), while LT within 30 days of listing was a positive predictor (35). NASH was not associated with survival differences in both studies (35,37).

Important considerations in the pre-transplant evaluation of patients with NASH and ACLF

Considering the strong evidence that NAFLD is associated with incident and prevalent hypertension and coronary artery disease and that it has high cardiovascular disease-related mortality (41), patients with NASH, who are diagnosed with ACLF should be considered early for transplant evaluation. Moreover, prompt cardiovascular assessment should be performed before these patients develop circulatory failure since this type of OF is more common in patients with NASH-related ACLF (23,24).

Therapeutic options

General principles of treatment of patients with NASH and ACLF

At present, there are no specific therapies for ACLF.

Table 2 Short-term mortality in two different populations of patients with ACLF and different etiologies of underlying liver disease and ACLF in patients with NAFLD

<table>
<thead>
<tr>
<th>Study</th>
<th>28-day mortality (%)</th>
<th>90-day mortality (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACLF entire cohort (CANONIC); Moreau et al. (7)</td>
<td>33.9</td>
<td>51.2</td>
</tr>
<tr>
<td>ACLF entire cohort; Hernaez et al. (21)</td>
<td>25.5</td>
<td>40.0</td>
</tr>
<tr>
<td>ACLF NAFLD based on EASL-CLIF criteria; Mahmud et al. (23)</td>
<td>45.2</td>
<td>59.2</td>
</tr>
<tr>
<td>ACLF NAFLD based on APASL criteria; Mahmud et al. (23)</td>
<td>51.0</td>
<td>73.5</td>
</tr>
</tbody>
</table>

ACLF, acute-on-chronic liver failure; NAFLD, non-alcoholic fatty liver disease; EASL-CLIF, the European Association for the Study of the Liver-Chronic Liver Failure; APASL, the Asian Pacific Association for the Study of the Liver.
The mainstay of treatment consists of identification and treatment of precipitating factors and management of OF (42). NASH patients should undergo prompt screening for infection and sepsis followed by immediate initiation of antibiotic therapy as infections were noted to be the most common precipitating event in this cohort (23,24). It is important when choosing empiric antibiotic therapy to take into consideration that nosocomial infections and infections caused by gram-positive bacteria or multi-drug resistant organisms are more frequent among ACLF patients (26). Fungal infections should be considered when there is no response to initial antibiotic therapy and/or the patients have concomitant T2DM. Adequate initial antibiotic therapy has been associated with better evolution and lower short-term mortality (26).

Role of artificial liver support systems in ACLF treatment

Artificial liver support systems are based on the concept of albumin dialysis and have been investigated as a potential option for bridging ACLF patients to LT. The best studied ones are the Molecular Adsorbent Recirculating System (MARS) and the Fractionated Plasma Separation and Adsorption system (Prometheus). The effect of these systems derives from their ability to remove albumin-bound toxins, nitric oxide, and pro-inflammatory cytokines (43). However, multicenter randomized controlled trials (RCTs) that investigated MARS vs. standard therapy (44) and Prometheus vs. standard therapy (45) failed to demonstrate benefit on 28- and 90-day survival. Another retrospective study showed MARS reduced early mortality at day 7 and 14 and suggested that the system might be more beneficial in patients with 2 or more OFs (46). The limitations of these studies were heterogeneity of patients’ selection and use of different ACLF definitions. A recent meta-analysis based on seven RCTs and 3 observational studies demonstrated a beneficial effect of artificial liver support systems on short-term survival in ACLF patients (47). Future studies of well-selected patients with ACLF should assess the utility of these support systems on survival of patients with NASH and assess whether their effect differs among patients with various etiologies of liver disease.

Role of granulocyte colony-stimulating factor (G-CSF) and cell therapy in ACLF

Other tested therapeutic options for ACLF are G-CSF administration and stem cell transplantation. The suggested beneficial effect of G-CSF is based on stem cell mobilization that could boost liver regeneration and on an immune modulation effect that can ameliorate immune dysfunction associated with ACLF. These potential effects may reduce the risk of infections (48). Two small RCTs (49,50) and a meta-analysis (51) have shown improvement in short-term survival, MELD, and Child-Pugh score, but due to the differences in ACLF definitions, population included, and variable doses and duration of G-CSF, it is difficult to make any firm conclusions on the efficacy of this treatment.

Animal data have demonstrated that transfusion of mesenchymal stem cells (MSCs) in hepatic failure may improve liver function, reduce inflammation, and reverse fibrosis (52). MSCs can be isolated from umbilical cord, bone marrow, or adipose tissue. The exact mechanism of MSCs for liver regeneration remains controversial, but it is considered they can differentiate into hepatocyte-like cells and integrate into the liver tissue, in addition to immunomodulatory effects and secretion of anti-inflammatory cytokines (53). To date, only one small open-label trial assessed the effect of umbilical cord-derived MSCs in patients with ACLF due to hepatitis B infection (54). It showed improved survival at 90 days and decreased MELD with no significant side effects. Currently, a phase 2 trial is investigating the safety of different doses of human liver-derived progenitor cells (HepaStem®, HHALPCs, Promethera Biosciences, Belgium) in patients with ACLF or with acute decompensation who are at risk for ACLF (NCT02946554). Another ongoing open-label trial has the aim to assess the safety and tolerability of HepaStem in cirrhotic and precirrhotic patients with NASH, but acute decompensation and ACLF are exclusion criteria (NCT03963921). The results of these trials are greatly awaited.

Conclusions

Despite the growing burden on NASH worldwide, there is a paucity of data that have assessed ACLF progression and outcomes in this population. Future studies should focus on the pathophysiology, clinical course, optimal LT evaluation, the impact of metabolic and cardiovascular comorbidities on LT candidacy and post-LT outcomes in NASH patients who develop ACLF.

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Footnote

Conflicts of Interest: The authors have no conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

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