Clinical Classification of Liver Failure: Consensus, Contradictions and New Recommendations

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Abstract

Liver failure (LF) is a clinical syndrome with complex clinical manifestations. The clinical diagnosis and classification of LF are still considerably different internationally. Based on the pace of the disease progression and its possible reversibility, LF can be divided into two categories: acute and chronic LF. However, a great difference exists in the diagnostic criteria between China, Europe, and the United States. These differences might be related to the various causes of LF, the complexity of clinical manifestations, and different experiences of experts from different centers. The main source of differences lies in the varied understanding of LF diagnostic standards.

The pathologic basis of various types of LF is different also. This article discusses the clinical diagnosis and classification of LF internationally and suggests new recommendations for classification of LF. According to this classification, ALF is divided into fulminant type and sub-acute type, and CLF is divided into acute exacerbative type and slowly progressive type. This classification, if adopted worldwide, could help achieve uniformity in the classification and therapeutic guidelines for liver failure. However, the validity of the classification should be further tested in clinical practice.

Keywords: Clinical manifestations; Chronic LF; Acute necrosis; Jaundice; Liver transplantation

Introduction

Liver failure (LF) is a clinical syndrome characterized by jaundice, coagulopathy, ascites, and hepatic encephalopathy. It is a devastating illness, with extremely high morbidity and mortality rates. Traditionally, LF is classified clinically as acute liver failure (ALF) or chronic liver failure (CLF). More recently, the entity of acute-on-chronic liver failure (ACLF) has been delineated. Pathological changes leading to LF consist of two types: 1) severe acute necrosis of liver tissues and 2) chronic progressive damage to liver cells [1]. The major causes of ALF and CLF are different, which can lead to different clinical manifestations and classification of LF in eastern and western countries: alcohol and drug abuse are the major causes of LF in the West, whereas viral infections are the predominant cause in the East [2].

In recent years, international organizations or societies have issued several guidelines or consensuses about the diagnosis and classification of LF [3-5]. In China, Guidelines for Liver Failure Diagnosis and Treatment were released in 2006 and updated in 2012 by the Liver Failure and Artificial Liver Group of the Chinese association of Infectious Diseases, and the Severe Liver Disease and Artificial Liver Group of the Chinese association of Hepatology [6,7]. These guidelines and consensuses have had a positive effect on LF-related research and academic exchanges worldwide. However, the clinical diagnosis and classification of LF still varies considerably among nations [8,9] and need further clarification.

The Connotation and Pathology of Liver Failure

The connotation of liver failure

Failure of the liver, the center of the body’s metabolism, is not limited to the liver itself, but also has a wide effect on the brain, kidneys, lungs, and other organs. Thus, LF may be considered multi-organ failure. First, we must recognize that LF is a functional diagnosis rather than a disease diagnosis. Second, we should focus on the difference between liver dysfunction and LF. Because the liver has a large reserve capacity and the ability to regenerate, mild or moderate liver damage usually does not result in overt dysfunction.

However, if the damage is more severe and widespread, as for example, from repeated or long-term injury, the resulting metabolic derangements can significantly affect hepatic function. These effects may consist of decreased detoxification of toxic substances, disorders of the formation and excretion of bile, and bleeding tendencies; these in turn may be manifested as jaundice, bleeding, or ascites, and other symptoms or signs. LF is a late stage of liver dysfunction, which may be expressed as hepatic encephalopathy, hepatorenal syndrome, and other disorders [10].
The pathology and clinical significance of liver failure

Studies suggest that pathological changes resulting in LF are of two main types: 1) Acute, severe liver necrosis and 2) chronic, progressive liver-cell damage [1]. There are some differences in pathological changes of LF, especially of ACLF. Some scholars have reported that the pathological changes associated with ACLF may be diverse and dominated by massive or sub-massive necrosis; cirrhosis accompanied with hepatitis may be one expression [11,12]. Li et al. [13] found that sub-massive hepatic necrosis is a pathological feature of hepatitis B virus (HBV)-associated ACLF.

Zhang et al. [14] found that pathological changes associated with ACLF are closely related to duration of disease. In decompensated cirrhosis, 70% of the liver cell function has been lost and is considered CLF [15]; it does not appear as areas or subareas of tissue necrosis. From a prognostic point of view, the significance of the distinction between ACLF and CLF lies mainly with patients who have a component of acute LF that could likely be reversed through liver regeneration. CLF usually is irreversible and may require liver transplantation for effective treatment [16].

Contradiction of Diagnosis and Classification of Liver Failure

Inconsistent LF classification

Although LF has been divided into acute and chronic LF, its classification is dissimilar in various countries and regions of the world. Guidelines and consensus for types of LF come mainly from Europe and the United States: they include the criteria for ALF recommended by the American association for Study of Liver Diseases (AASLD), in 2005 [2]; ACLF consensus released by the Asia-Pacific Association for the study of Liver (APASL), in 2009 [16]; and the consensus on ACLF by AASLD and the European association for the Study of Liver (EASL), in 2011 [4].

The CLF guidelines are non-uniform, only appropriate guidelines or consensus for the complications of cirrhosis, such as ascites, hepatic encephalopathy, and hepatorenal syndrome, have been published [17-19].

Chronic Liver Failure, Mechanisms and Management, edited by Gines et al., were published in 2011 [2]. Williams [20] proposed that LF be divided into three categories: ALF, ACLF, and CLF. According to the Liver Failure Guidelines in China, issued in 2006 and revised in 2012, LF is divided into four categories: ALF, sub-acute LF (SALF), ACLF and CLF (Table 1).

The establishment of CLF as a distinct entity is necessary in order to maintain the continuity and integrity of LF classification. However, there is lack of uniformity of CLF guidelines. The cause of this lack might be thought of as the decompensated stage of liver cirrhosis, and appropriate guidelines or consensus for complications of cirrhosis are those mentioned above.

Table 1: The current main categories of liver failure and definitions.

<table>
<thead>
<tr>
<th>Three categories (United Kingdom) [20]: ALF, ACLF and CLF</th>
<th>Four categories (China) [7]: ALF, SALF, ACLF and CLF</th>
</tr>
</thead>
<tbody>
<tr>
<td>Definitions</td>
<td>ALF defined as HE within 6 weeks</td>
</tr>
<tr>
<td>SALF defined as clinical manifestations of liver failure within a period of 15 days-25 weeks.</td>
<td></td>
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<tr>
<td>ACLF defined as acute deterioration of preexisting chronic liver disease, usually related to sepsis, alcohol, or bleeding.</td>
<td></td>
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<tr>
<td>CLF defined as progression of end-stage liver disease.</td>
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<tr>
<td>CLF defined as chronic deterioration of preexisting liver cirrhosis.</td>
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</tr>
</tbody>
</table>

ALF: Acute Liver Failure; SALF: Sub-acute Liver Failure; ACLF: Acute-On-Chronic Liver Failure; CLF: Chronic Liver Failure; HE: Hepatic Encephalopathy; TBIL: Total Bilirubin; PTA: Prothrombin Activity.

Diagnostic Criteria Differences for Various Types of LF

Given the lack of generally accepted, evidence-based diagnostic criteria, the diagnosis of LF varies within China and between China and other countries. These differences might be due to the variety of causes of LF, the complexity of its clinical manifestations, and differences in expert opinions from various centers. For example, alcoholic cirrhosis constitutes 50% to 70% of all underlying liver diseases of ACLF in western countries, whereas hepatitis B- or C-related cirrhosis constitutes about 10% to 30%. In most Asian countries, however, hepatitis B constitutes about 70% of ACLF, and alcohol abuse only approximately 15% [21]. These factors notwithstanding, the more important barrier to achieving consensus in LF classification lies in the variations in LF diagnostic standards. For example, the current definition of ACLF differs greatly in various countries. The APASL definition stresses the occurrence of ascites and/or encephalopathy occurring within a period of four weeks in patients with underlying chronic liver disease, whereas the AASLD/EASL definition underlines the occurrence of multi-organ failure in patients with chronic liver disease, resulting in three-month mortality (Table 2). This difference has led to a misconception between ACLF and acute decompensation of liver cirrhosis [22,23]. Moreau et al. [24] reported that ACLF is a different syndrome from that of acute decompensation of cirrhosis. Since the majority of ACLF patients in the Moreau study had alcohol-induced cirrhosis, this conclusion cannot be extended to virus-related ACLF [25]. Because the diagnostic criteria differ significantly, the prognoses of ACLF patients may differ. Thus, Zhang et al. [26] reported that the 90-day mortality rates in...
the ACLF diagnostic criteria established by APASL, CMA and EASL are 13.1%, 25.3% and 59.3%, respectively. Wang et al. [27] also indicated that the diagnostic criteria for ACLF established by EASL are stringent and may impede early therapeutic intervention. In regard to ALF, Wlodzimirow et al. [28] reported that 41 different definitions of ALF were used in 87 studies. The diversity in definitions of ALF hinders the making of comparisons and quantitative analyses among studies. Whereas there is room for improvement in the reporting of ALF definitions in prognostic studies [28], scholars generally agree on the definition of CLF [6,20].

Table 2: Definitions of ALF and ACLF by major societies.

<table>
<thead>
<tr>
<th>LF</th>
<th>Proposed by</th>
<th>Definitions</th>
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<tbody>
<tr>
<td>ALF</td>
<td>AASLD (2006)</td>
<td>No previous history of cirrhosis; deterioration of liver function occurred in 26 weeks; coagulopathy (INR ≥ 1.5); any degree of altered consciousness (encephalopathy).</td>
</tr>
<tr>
<td>ACLF</td>
<td>APASL (2009)</td>
<td>Acute hepatic insult manifested as jaundice and coagulopathy, complicated within 4 weeks by ascites and/or encephalopathy in a patient with previously diagnosed or undiagnosed chronic liver disease.</td>
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<td></td>
<td>AASLD/EASL (2011)</td>
<td>Acute deterioration of preexisting chronic liver disease usually related to a precipitating event and associated with increased mortality at 3 months due to multisystem organ failure.</td>
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<td></td>
<td>EASL-CLIF Consortium (2013)</td>
<td>Acute liver deterioration, organ failure (defined by the chronic liver failure-sequential organ failure assessment score) and high 28-day mortality rate (&gt; 15%) on the basis of acute decompensation of liver cirrhosis.</td>
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<td></td>
<td>WGO working party (2014)</td>
<td>ACLF may be divided into 3 categories depending on whether or not there is underlying cirrhosis and hepatic decompensation: Type A, non-cirrhotic ACLF; Type B, cirrhotic ACLF; Type C, cirrhotic ACLF with previous hepatic decompensation.</td>
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Proposed New Classification and Diagnosis of Liver Failure

Given the disagreements of LF diagnosis and classification, the author believes that the diagnosis of hepatic failure must be separated from the clinical classification of complications caused by LF, such as hepatorenal syndrome and hepatic encephalopathy. Although the complications caused by LF are different and have inconsistent manifestations at different stages, the final clinical manifestations are essentially the same. The different stages of the disease are the most significant factors for the prognosis of the patient. Therefore, relaxation of the diagnostic criteria for LF should emphasize prevention in favor of improving patients’ prognosis [33]. For assessing efficacy of treatment, standards should be uniform internationally. Thus, we propose that, based on ALF and CLF, the former be further divided into fulminant-type and subacute-type hepatic failure, and the latter be divided into acute exacerbative type and slowly progressive type (Figure 1) [34]. These criteria of LF are different from the current criteria of the EASL, AASLD and APASL, which might be simpler and more practical.

Different Pathology Bases of Various Types of LF

Significant differences in understanding of ACLF still exist locally and internationally. Most scholars in Europe and the United States believe that the basic underlying disease is compensated cirrhosis [4,29]. Moreau et al. [24] have stated that decompensated cirrhosis is the basis of ACLF. In China, the ACLF diagnostic criteria take into account the degree of chronic liver disease (e.g., chronic hepatitis and liver cirrhosis), which causes much confusion. ACLF patients have high short-term mortality. However, those who survive the acute exacerbation live longer than do patients with decompensated cirrhosis [30]. Recently, the World Gastroenterology Organization Working Party suggested that ACLF be divided into 3 categories depending on whether there is underlying cirrhosis and hepatic decompensation: Type A, non-cirrhotic ACLF; Type B, cirrhotic ACLF; Type C, cirrhotic ACLF with previous hepatic decompensation [31]. According to the diagnostic criteria of ALF or subacute LF in China, these types should know the basis of chronic liver disease and are different from ALF as defined by the AASLD [2]. ALF patients have no previous history of cirrhosis, and the deterioration of the liver function occurred within 26 weeks according to the AASLD definition. Also, ALF is characterized by coagulopathy (prothrombin international normalized ratio ≥ 1.5) and any degree of altered consciousness (encephalopathy). For Wilson’s disease, vertical transmission of hepatitis B, or autoimmune hepatitis, disease found at 26 weeks may also be considered ALF even in the presence of cirrhosis. Considering that most Chinese chronic hepatitis B patients are long-term carriers of hepatitis B virus, the first severe occurrence of CHB were regarded as ACLF [32]. Therefore, the current ACLF diagnostic criteria in China might include some ALF due to acute or subacute hepatic necrosis.
jaundice, ascites, and bleeding tendencies, but no hepatic encephalopathy; although their disease progresses relatively slowly, their prognosis is poor (the mortality rate more than 40%~50%). We believe that this condition should be designated subacute LF [37]. Accordingly, ALF could be divided into fulminant and sub-acute types, with the latter having a four-week duration of disease. The designation of fulminant type requires hepatic encephalopathy, whereas the sub-acute type does not necessarily have hepatic encephalopathy and is mainly characterized by severe jaundice and ascites etc.

**CLF**

Patients with decompensated cirrhosis, because of complications or other factors, may experience either acute deterioration of liver function or slowly progressive disease. Acute deterioration of liver function is defined as the TBil ≥ 171 umol/L and PTA ≥ 40%. Accordingly, CLF can be divided into two types: slowly progressive and acute exacerbarative type. The slowly progressive type is equivalent to the current four-week deterioration of liver function in patients with hepatic encephalopathy. Acute exacerbarative type is equivalent to deterioration that occurs in decompensated cirrhosis. The short-term prognosis of this type was worse than that of the slowly progressive type [35,38]. Acute exacerbarative type might include the ACLF proposed by EASL, which is more serious yet [26].

**Conclusion**

In summary, considering the numerous causes of LF and clinical manifestations caused by liver damage, differences in the classification of LF are to be expected. Some definitions of LF as ACLF might be conducive to evaluation of short-term prognosis and be harmful to early intervention of disease, versus other definitions. This classification of LF proposed by the author might be more suitable for determining therapy and estimating prognosis. The connotation of ALF and CLF in this classification are different from that of the EASL, AASLD, APASL and China although the name of ALF and CLF are as same as the previous reports [3-7]. Hence, our classifications are new and more rational classification scheme. This classification, if adopted worldwide, could help achieve uniformity in the classification and therapeutic guidelines for liver failure. However, the validity of this classification should be further tested in clinical practice.

**References**


