Liver Failure due to Acute Viral Hepatitis (A–E)

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Introduction

Acute liver failure (ALF) is a devastating clinical syndrome associated with high mortality in the absence of immediate state-of-the-art intensive care, specific treatment, or liver transplantation. It is partly accompanied by the onset of hepatic encephalopathy within 8 weeks of the first symptoms [1]. The American Association for the Study of Liver Disease (AASLD) position paper recommends that all patients with clinical or laboratory evidence of acute hepatitis should have immediate measurement of prothrombin time and careful evaluation for subtle alterations in mentation [2]. The most widely accepted definition proposed by the AASLD includes evidence of coagulopathy (international normalized ratio $\geq 1.5$) and presence of an altered sensorium (encephalopathy) without pre-existing cirrhosis and with duration of symptoms of less than 26 weeks [3]. For patients meeting this definition, hospital admission, ideally to a transplant center, is mandatory [2].

ALF can occur as a result of various etiologies (overdosing with acetaminophen or other drugs, viral hepatitis, ischemia, and other causes). When ALF is not fatal, the liver has a unique capacity to recover completely, although reliably predicting mortality remains challenging [4, 5].

Causes and Epidemiology

ALF is a rare disease especially in the developed world [3]. The causes of ALF are heterogeneous and may vary from country to country. Traditionally, hepatitis B was regarded as the most common cause of ALF [6]. Over the last decade, statistics suggest a change in ALF etiology in developed countries. According to the United States ALF Group Registry statistics, drug-induced liver injury (DILI), including acetaminophen and idiosyncratic related etiologies, is responsible for more than 50% of cases, followed by in-
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Fig. 1. Causes of ALF worldwide. *Others include non-A–E viral hepatitis; **no discrimination between acetaminophen and idiosyncratic drug injury [3, 11, 17, 71–76].

Determinable causes (14%), hepatitis B (7.7%), and autoimmune hepatitis (5.9%), followed by less common causes including ischemia, Wilson's disease, Budd-Chiari syndrome, and pregnancy-related liver failure (e.g. HELLP syndrome) [7, 8]. Statistics from Northern European countries and the UK stand in line with statistics from the USA [3, 9]. Reasons for the decline in viral-related cases of ALF range from successful vaccination programs over improved control of blood products to a more health-educated society [10]. However, statistics from Southern Europe are contrary where viral hepatitis still is a main cause of ALF [11, 12]. Therefore we can assume a north-south gradient within Europe. Predictions for Eastern Europe will become more sophisticated as statistics with reasonable numbers of patients are reported. As most of those infected with hepatitis B virus (HBV) in the European region live in Eastern European countries [13, 14], a severe impact in the role of ALF is likely.

Bernal and Wendon [3] assume that, globally, hepatitis E and A infections are probably responsible for the majority of ALF cases. This can be related to high incidence in some Asian countries. In India, for example, one study reported that 44% of ALF cases were related to hepatitis E virus (HEV) infection. Similar results were obtained in Bangladesh [15, 16].

In Germany, findings can be categorized alongside recent developments in Europe. Drug toxicity appears to be the major cause of primary ALF. In contrast to findings in the USA and the UK, idiosyncratic drug injuries, rather than acetaminophen, are most commonly related to DILI. Additionally, hepatitis B, with 18% of all cases, still plays a major role in etiology. The substantial proportion of unknown cases and the fact that the diagnosis of idiosyncratic liver injury is based on the exclusion of other diagnoses leaves the question open whether other etiologies might be overlooked [17]. Indeed, we could prove recently that a reasonable number of patients with ALF might be related to unappreciated hepatitis E infections. Figure 1 provides an overview of the causes of ALF worldwide [18].

Hepatotropic Viruses

**Hepatitis A**

Hepatitis A virus (HAV) can cause liver disease that ranges from mild to severe illness but does not cause chronic liver disease. It is usually transmitted through ingestion of contaminated food and water or through direct contact with infectious persons. Thus, the risk is higher in areas with a lack of safe water and poor sanitation. Transmission by blood transfusion is rare; however, polymerase chain reaction (PCR) testing for blood donors is recommended. Measures to improve these issues as well as vaccination are known...
to be most effective to combat the disease. Usually most patients make a full recovery from hepatitis A [19]. Unfortunately, a small proportion of infected patients develop a fulminant hepatitis with a high mortality rate. The World Health Organization (WHO) calculates that 1.4 million people become infected with HAV each year [20, 21].

In Germany, 681 cases of hepatitis A were reported to the Robert Koch Institute (RKI) in 2014. This is the lowest rate since 2001, and the prevalence dropped to 0.8/100,000 inhabitants. Most of the patients acquired their infection within Germany (63%) [22].

Less than 1% of acute HAV infections result in ALF [20]. Young children generally belong to the group of patients with unapparent or subclinical hepatitis and have no symptoms or jaundice [23]. In contrast, the infection is more severe in adults, with symptoms occurring in 70%. Generally, hepatitis A-related ALF has a spontaneous survival rate of 69%; the remaining 31% require emergency liver transplant (ELT) or die [24]. Furthermore, patients with pre-existing liver damage such as non-alcoholic fatty liver disease (NAFLD) or alcoholic steatohepatitis (ASH) are more susceptible to develop an acute-on-chronic liver failure in cases of HAV infection [25].

The reasons why HAV infection may progress infrequently to ALF are poorly understood. Underlying host factors such as age and even minor pre-existing liver damage may play a role [26, 27]. Moreover, viral factors including low viral load and a higher rate of substitutions in the 5′ untranslated region of the viral genome are also thought to increase the likelihood of a fulminant course [28, 29]. Investigations to find a clear difference in genomic sequences between patients with fulminant courses of HAV and those with a minor course did not lead to clear results [30]. In contrast, there is some evidence that cytolytic T cells may play an important role in the pathogenesis of HAV infection and in determining the course [31]. The declining incidence of HAV infection in developed countries makes it unlikely that great efforts will be taken to target this problem in the near future.

Specific diagnosis can be made by the detection of HAV-specific immunoglobulin (Ig) M and IgG antibodies in serum. Additional tests may include reverse transcription PCR (RT-PCR) to detect the viral RNA. There is currently no specific treatment. With acute hepatitis A, as with many other etiologies of ALF, care is mainly supportive (according to the AASLD guidelines), and if recovery is unlikely, liver transplantation should be considered [2]. Data for outcomes after transplantation is not available.

Hepatitis B

Hepatitis B is a viral infection caused by the HBV and can lead to both acute and chronic liver disease. HBV is a DNA virus and a member of the Hepadnaviridae family that is transmitted through contact with blood or other body fluids of an infected person. In highly endemic areas, hepatitis B is most commonly spread from mother to child at birth (perinatal transmission). In industrialized countries, hepatitis B is mostly spread by sexual transmission. Other modes of transmission include the reuse of needles and syringes either in healthcare settings or among persons who inject drugs; medical, surgical, and dental procedures; tattooing; and the use of razors and similar objects that are contaminated with infected blood [32, 33].

In Germany, the hepatitis B surface antigen (HBsAg) prevalence is less than 1%. In 2014, the RKI reported 755 new infections (0.9 cases per 100,000 inhabitants). A rise in infection rates has been especially observed in metropolitan regions. However, rates in some Asian, African, and Eastern European parts are much higher [22].

In adults, hepatitis B infection is usually asymptomatic. Young children are most likely to develop a chronic infection whereas in adults, less than 5% of otherwise healthy persons who get infected will develop chronic infection [34]. According to the WHO, one quarter of the world’s population (2 billion) is estimated to have experienced HBV infection and 3% (240 million) are chronically infected with hepatitis B [35]. About 1% of persons with acute hepatitis develop ALF, which can be regarded as a more atypical course of acute hepatitis [36]. However, in a fulminant course, HBV DNA and HBsAg levels often fall rapidly as liver failure develops; thus, some patients are HBsAg-negative by the time of onset of hepatic coma [37]. Therefore, a transfer to a specialized unit and accurate testing are crucial (table 1). Reasons why some patients develop a fulminant course in comparison to others are not well investigated. Simultaneous intake of alcohol, acetaminophen, or methamphetamine may play a role [38]. In addition, HBV genotype might affect the outcome. In one study comparing the genotypes of chronic HBV patients to cases with HBV-related ALF, the latter showed a higher prevalence of genotype D [39].

The mechanisms whereby HBV induces ALF are not well known. There are several studies showing that HBV core promoter mutations have been implicated in the pathogenesis of fulminant hepatitis B. A common hallmark of these mutations is a phenotype of enhanced viral replication. It is common in patients with a more aggressive course such as fulminant hepatitis but also in chronic hepatitis. Enhanced viral replication may result in a stronger immune response and extended infection. Consequently, this could be an important factor in driving the disease towards a fulminant course. Moreover, there is also evidence that at least some mutated variants may induce apoptosis in hepatocytes [40, 41]. Furthermore, it is worthwhile mentioning that patients with underlying chronic liver disease are more susceptible to develop acute-on-chronic liver failure if infected with certain mutants [42]. However, since patients with fulminant hepatitis exhibit different phenotypes be an important factor in driving the disease towards a fulminant course. Moreover, there is also evidence that at least some mutated variants may induce apoptosis in hepatocytes [40, 41]. Furthermore, it is worthwhile mentioning that patients with underlying chronic liver disease are more susceptible to develop acute-on-chronic liver failure if infected with certain mutants [42]. However, since patients with fulminant hepatitis exhibit different phenotypes so that a highly replicative phenotype is not exclusive, additional mechanisms must be involved.

| Table 1. Initial laboratory tests to perform in the initial evaluation of ALF (HAV, HBV, HEV) [2, 18] |
|-------------------------------|-----------------------------------|-------------------------------|--------------------------|
| HAV IgM antibody to HAV (IgM anti-HAV) | HBV hepatitis B surface antigen (HBsAg) | HBV IgM antibody to hepatitis B core (IgM anti-HBc) | HEV HEV RNA PCR |
| HAV IgG and IgM antibody to HAV (IgG/IgM anti-HAV) | HBV DNA PCR | HEV IgG and IgM antibody to HEV |  |
Another possible factor with major impact in HBV-related ALF could be the overwhelming activation of humoral immunity. A massive accumulation of plasma cells secreting IgG and IgM and complement components were found in necrotic areas in livers of HBV-related ALF. Furthermore, these antibodies were found to target hepatitis B core antigen (HBcAg) [43]. How relevant these results are for HBV-related ALF and underlying mechanisms needs further investigation, as this particular study has been performed with only 2 patients.

The incidence of reactivation of previously stable subclinical infection with HBV without a manifestation of chronic disease characteristics is steadily rising. This is related to patients with treatment-induced immunosuppression during or after cancer therapy. Therefore, identification of patients at high risk as well as the use of prophylaxis is crucial [44]. High disease activity usually leads to clinical and serological resolution. However, even after serological resolution, small amounts of covalently closed circular DNA (ccc-DNA) persist in the liver for years, decades, and possibly for the lifetime. T cell immunity suppresses viral replication from these ccc-DNA copies to very low levels [45]. Mutations in the HBV genome, immunosuppressive therapy, and viral- or drug-induced injury are common causes of reactivation. Fulminant hepatic failure in the setting of HBV reactivation has a much higher morbidity and mortality rate than de novo fulminant hepatitis [46]. The AASLD recommends that patients found to be positive for HBsAg who are to begin such a therapy should be treated prophylactically with a nucleoside analog and that it should be continued for 6 months after completion of immunosuppressive therapy. Moreover, the American Gastroenterological Association recommends in their guidelines to offer HBsAg-negative but anti-HBc-positive patients a prophylactic therapy if treated with B-cell-depleting agents (eg, rituximab) [47].

The prognosis of HBV-related ALF is generally limited without liver transplantation. Transplant-free survival rates range from 26 to 53% [48]. Hepatitis serological testing should be done for identification of acute viral infection. This should also be performed in case another putative etiology has been identified [2].

To date, there is not enough evidence that artificial liver support devices can enhance the patient’s survival. In contrast, there is first evidence that initial antiviral therapy could be beneficial. Jochum et al. [49] could show that immediate treatment of HBV-induced ALF with drugs such as entecavir is well tolerated and beneficially affects the course of the disease. Therefore, the AASLD recommends to take nucleos(t)ide analogs into consideration as a possible treatment option for hepatitis B ALF as well as for the prevention of post-transplant recurrence [2].

**Hepatitis C**

Although controversial, currently there is no evidence that hepatitis C alone appears to cause ALF. However, an acute hepatitis C is able to induce hepatic damage and a severe increase of transaminases. Nevertheless, this does happen without altered synthesis function [50].

**Hepatitis D**

Hepatitis D virus (HDV) can be regarded as an altered RNA virus as it is dependent on the coexistence of HBV. The virus consists of only one protein, a circular RNA strand, i.e. the actual hepatitis D antigen. It requires the HBsAg for the encapsidation of its own genome. It is further dependent on the sharing of envelope proteins to facilitate the assembly of the HDV genome into new virus particles. Apart from that, the replication of the circular RNA is independent of HBV [51, 52].

In Germany, 17 cases of hepatitis D were reported to the RKI in 2014 [22]. Hepatitis D coinfection may occur in conjunction with acute hepatitis B. In this case, the rate of hepatitis B to progress to ALF approaches 20%. In addition, HDV superinfection of a chronic HBV patient can also result in ALF [53]. Although there is no specific therapy for hepatitis D infection, diagnosis is important, as the disease follows a characteristic biphasic pattern in which initial recovery is followed by clinical deterioration.

**Hepatitis E**

Approximately 20–40% of ALF cases are caused by HEV infections in developing countries. However, in western countries, liver failure caused by hepatitis E has only been recently regarded as an imported infection. In line with this opinion, data from surveys based on population from the USA showed hepatitis E as a rare cause of ALF [54]. However, recent studies showed HEV seroprevalence rates between 16 and 20% among adults in industrialized countries [55–57]. Nevertheless, these numbers should be interpreted with caution as results may be influenced by serological assay discrepancies. Despite these inter-assay variations, there is some evidence of autochthonous HEV, particularly genotype 3, being endemic to Europe [58, 59].

In contrast to ALF, a study by Davern et al. [60] from 2013 showed that in a US multicenter study 9 (3%) patients of 318 cases of suspected DILI were misdiagnosed. After reassessment these cases were considered more likely to be hepatitis E-related [60]. This study is supported by another study, which showed that 22.2% of patients diagnosed with autochthonous hepatitis E were initially thought to have DILI prior to HEV testing. Moreover, the investigators could demonstrate that 21.4% of patients with criterion-referenced DILI had autochthonous hepatitis E after the reassessment of stored serum with HEV testing [61]. These studies underline that a structured assessment of medical history can be difficult. Therefore, it is necessary for western clinicians to take HEV into account as a cause of liver disease when making their treatment assessments.

Indeed, this should also be the case in the setting of ALF. Contrary to results from the USA, one retrospective study from Germany suggests that hepatitis E contributes to a substantial proportion of acute liver injury cases in developed countries [60, 62]. In a retrospective analysis of 80 ALF cases, 8 of these cases have been detected to be potentially hepatitis E-related, suggesting that HEV testing is necessary in every case of ALF. Moreover, this study suggests that it is necessary to perform both serological testing and RNA testing as well [18].
In immunocompetent patients, HEV infection usually takes an asymptomatic course [63]. In contrast to descriptions of a higher incidence of chronic hepatitis E in immunocompromised patients, two case reports of acute hepatitis E claim that steroids may prevent the progression of acute hepatitis E during ALF [64, 65]. Further studies are needed to evaluate whether steroid treatment has a supportive role in hepatitis E-related acute liver injury. However, patients with underlying liver disease as well as the elderly are more likely to show a more dramatic ALF course [62].

There is little known about the impact of acute infections and the likelihood of exacerbation to fulminant liver injury in the setting of immunosuppression and hematopoietic stem cell transplantation (HSCT). Recently, a fatal outcome of autochthonous hepatitis E in a patient with B-cell lymphoma in France has been reported [66]. However, contrary to HBV, descriptions of fulminant courses in hematological malignancies are rare.

Considering all of the above, the question arises whether antiviral therapies might be effective against HEV-related ALF. There is only little evidence for the use of pegylated interferon alpha and ribavirin to treat chronic HEV infection [67–69]. Ribavirin also seems to be a possible treatment option in acute hepatitis [67]. Descriptions of treatment in ALF are rare although there are occasional reports of successful treatment in acute-on-chronic liver failure [70]. Further trials must be conducted to clarify the need and benefit of any antiviral treatment.

Disclosure Statement

The authors have nothing to disclose.

References

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