Abnormal Liver Blood Tests: Hepatologist Approach

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Abstract
Background: Available data suggest that the prevalence of chronic liver disease (CLD) and primary liver cancer is rising in Europe and represents a major public health problem. Predictions are showing that these trends will continue to rise in the upcoming years. Summary: Alcohol-related liver disease, nonalcohol fatty liver disease, and viral hepatitis B and hepatitis C are the leading causes of liver cirrhosis and primary liver cancer in Europe. Drug-induced liver injury represents a major cause of acute hepatitis, while liver transplantation is the second most common solid organ transplantation in the world. Patients with CLD have increasing rates of hospitalization, longer hospital stays, and more adverse outcomes compared to the other chronic conditions. Direct targeting of risk factors can prevent complications of advanced liver disease and improve outcome. Patients with CLD should be referred to a hepatologist for assessment of the stage of liver disease, for specific treatment and screening for hepatocellular carcinoma. Moreover, patients with unknown etiology of abnormal liver blood tests should be referred to a hepatologist for assessment of liver disease, as well as for prevention and treatment of complications of cirrhosis and/or portal hypertension. Key Messages: CLD is amenable to prevention and treatment, while disease management strategies need to improve in order to reduce the burden of liver disease and deaths due to end-stage liver diseases.

Introduction
Chronic liver disease (CLD) represents an area of clinical priority due to increasing rate of morbidity and mortality. Cirrhosis is the 11th leading cause of death worldwide, while liver disease accounts for 2 million deaths per year globally [1, 2]. Approximately 14% of US adults are diagnosed with alcohol use disorders and are at high risk for alcohol-related liver disease (ARLD). Approximately 2 billion adults are obese worldwide, and over 400 million adults have diabetes, both of which are risk factors for nonalcoholic fatty liver disease (NAFLD) [2]. Drug-in-
duced liver injury (DILI) represents a major cause of acute hepatitis, while the prevalence of viral hepatitis remains high worldwide. Less common causes of CLD are primary biliary cholangitis, autoimmune hepatitis (AIH), and primary sclerosing cholangitis. Hereditary hemochromatosis is a common genetic disorder affecting 1 in 200 people in Europe and USA, while rare genetic disorders are Wilson’s disease and alpha-1 antitrypsin deficiency. Liver cancer is the 16th most common cause of death. Liver transplantation is the second most common solid organ transplantation in the world [2]. Predictions are showing that trends will continue to rise in the upcoming years, predominantly from increasing rate of NAFLD and ARLD [3, 4]. Patients with CLD in comparison with other chronic disorders have increased rates of hospitalization, longer hospital stay, more complications, and more adverse outcomes in general [3]. Disease management models for CLD greatly need to manage the anticipated increase in burden of liver disease and hospitalizations for CLD.

**Nonalcoholic Fatty Liver Disease**

NAFLD is the most frequent liver disease in Western Europe and USA, with the prevalence ranging from 17 to 46% of adults. NAFLD is defined as hepatic fat accumulation in >5% of hepatocytes according to histology and can be presented with a spectrum of disease activity from NAFL, early NASH, fibrotic NASH, cirrhosis, and HCC [4].

Patients who have metabolic risk factors (waist circumference 94/80 cm for men/women and/or BMI >25), arterial hypertension, dyslipidemia, insulin resistance, or diabetes type 2 (T2DM) should be further assessed for NAFLD [5]. NASH and fibrotic NASH are rapidly progressive diseases and are associated with an increased mortality rate in comparison with NAFL. Emerging body of evidence suggests that NAFLD is a multisystemic disease associated with cardiovascular disease (CVD), CKD, T2DM, and reduced mineral density [6]. The most common cause of death in patients with NAFLD is CVD, independent of other metabolic comorbidities [5]. At present, NAFLD with advanced liver fibrosis is the third most common cause of HCC and is considered the second leading indication for HCC-related liver transplantation (LT) [7].

NAFLD is present in 7% of nonobese individuals (lean NAFLD) [8]. Several genetic polymorphisms on the *PNPLA3* gene and *TM6SF2* gene have been identified with associated risks for NAFLD and increased risk for HCC, but genotyping is not recommended routinely in clinical practice, so far [9, 10].

Patients with NAFLD should be referred for noninvasive diagnostic algorithm to predict NASH and fibrosis and should be evaluated for NAFLD-associated diseases (T2DM, CVD, and dyslipidemia). Patients with inconclusive tests should be referred to a hepatologist for transient elastography. In patients with NASH fibrosis, the final diagnosis should be made by liver biopsy [4, 6].

Individuals with NAFLD require reinforcement of lifestyle changes regarding dietary restriction and increase in physical activity with aerobic exercise in combination with resistance training. Pharmacologic therapy should be reserved only for patients with progressive NASH, active NASH, and for early-stage NASH with increased risk of fibrosis (age >50 years, T2DM, increased ALT, and MetS) [4]. Current therapy includes pioglitazone, vitamin E, or their combination. Statins can be used to prevent cardiovascular risk, but with no effects on liver disease [11].

**Alcohol-Related Liver Disease**

According to the WHO, the highest alcohol-related morbidity and mortality is in Europe, with the mean alcohol consumption of 10.9 L of pure alcohol per person per year. In the EU, 41% of the liver deaths are attributed to alcohol. Liver injury due to alcohol depends on several more factors including types of alcohol, duration of exposure, genetic susceptibility, and drinking patterns [12, 13]. Alcohol screening questionnaires are very helpful and should be done systematically by physicians [14]. The AUDIT (Alcohol Use Disorders Inventory Test) remains the gold standard, as it has good sensitivity and specificity to identify heavy drinkers and explicitly addresses consumption dependence [15]. Shorter version, the AUDIT-C, is reliable for the screening of risky drinking occasions [16, 17]. The CAGE is an easy-to-use tool to identify severe alcohol dependence, although it appears to be a poor screening questionnaire for heavy drinking [18].

Diagnosis of ARLD can be established upon history of regular alcohol consumption (>20 g/day for women and >30 g/day for men), together with the presence of clinical, laboratory, and imaging techniques suggestive of liver injury (ultrasonography, CT, and MR) [13]. Alcoholic hepatitis (AH) is a distinct clinical syndrome characterized by the acute onset and progression of jaundice with or
without other signs of liver decompensation (encephalopathy and ascites) in patients with active alcohol abuse [13]. Diagnosis of AH is based on clinical and typical laboratory findings in a patient with a history of recent heavy alcohol use. Early improvements in liver function have a major impact on mortality rate in AH.

The Maddrey score is the most widely used prognostic model for assessment of risk of early death from AH in the short term [19]. More recently, several prognostic scores have been developed such as MELD (Model for End-Stage Liver Disease) score [20]. Another prognostic score, the Lile model, is based on the response of serum levels of bilirubin to a 7-day treatment of corticosteroid therapy [21, 22].

A liver biopsy is reserved for patients where there is diagnostic uncertainty, if there is suspected additional causes of liver injury, to establish the definite diagnosis, to assess the stage of the liver disease, and for prognosis. Approximately 10–20% of patients with ARLD have co-existing etiology of their liver disease [13].

Therapeutic measures include alcohol abstinence regardless of the stage of the liver disease, adequate nutrition support in malnourished patients (sarcopenia is present in 50% of patients with advanced ARLD), supplementation with vitamin B complex, treatment of ascites, hepatic encephalopathy (lactulose and rifaximin), and preventive measures of variceal bleeding [13, 23]. Coffee drinking seems to decrease the risk of progression to cirrhosis [24]. Treatment algorithm for patients with AH is shown in Figure 1.

**Hepatitis C Virus Infection**

Hepatitis C virus (HCV) presents one of the most common causes of CLD. It is estimated that 71 million individuals worldwide are infected with HCV [25]. According to the WHO, novel therapies and enhanced improvement in understanding the pathophysiology of HCV infection make a possible target to eradicate hepatitis C worldwide in the near future [26].

At least 8 major genotypes of HCV (each comprising multiple subtypes) have been identified worldwide. HCV genotypes 1, 2, and 3 appear to have a worldwide distribution, while the relative prevalence of all genotypes varies from one geographic region to another.
The diagnosis is based on presence of anti-HCV antibodies by enzyme immunoassay and detectable HCV RNA in serum by a sensitive molecular assay with a limit of detection <15 IU/mL. Anti-HCV antibodies may be undetectable in the early phase of acute infection and in individuals who are immunosuppressed. Severity of liver disease must be assessed prior to treatment. Identifying patients with cirrhosis or advanced fibrosis is important because the specific treatment choice depends on the stage of fibrosis [25].

The primary therapeutic goal is to achieve a sustained virologic response. It is defined as undetectable HCV RNA after treatment completion (in <0.2% of cases, relapse occurs). The combinations of HCV drugs available in Europe are as follows: sofosbuvir, sofosbuvir/velpatasvir, sofosbuvir/velpatasvir/voxilaprevir, glecaprevir/pi-brentasvir, and grazoprevir/elbasvir. Treatment with sofosbuvir/velpatasvir and glecaprevir/pi-brentasvir can be started without knowledge of the genotype or subtype with high probability of success. In geographical areas (Asia, South America, and Africa) where HCV subtypes inherently resistant to NS5A inhibitors are present, the HCV genotype and subtype should be determined with an assay that can discriminate subtype 1a from 1b, if possible [25].

All individuals with HCV infection, who are either treatment naïve or treatment experienced, should be treated without delay. Patients with decompensated cirrhosis with or without an indication for LT should be treated in order to achieve improvement in liver function and survival. For patients awaiting LT, the goal is not only to improve liver function, but also to prevent liver graft infection after LT [27, 28].

HCC incidence has dramatically decreased in patients with HCV or HBV infection due to highly effective antiviral therapy [25, 29]. However, patients with cirrhosis who eradicate their HCV infection remain at substantial risk of HCC. The risk increases with age, advanced chronic disease, cirrhosis, and presence of diabetes. Thus, patients with HCV-induced cirrhosis should be included in follow-up for early detection of liver cancer [29, 30].

**Hepatitis B Virus Infection**

Hepatitis B virus (HBV) infection remains still an important global health problem with approximately 240 million people infected worldwide [31]. The highest prevalence is in sub-Saharan Africa and East Asia, where 5–10% of adult people are chronically infected. Vertical transmission from the infected mother to child – perinatal transmission – leads to chronic disease in >90% of newborns in these areas [31, 32]. On the other hand, the prevalence of HBV infection is constantly decreasing due to implementation of the vaccination programs worldwide, improvements in the socioeconomic status, and possibly due to effective antiviral treatments [33].

Patients with HBV infection should be assessed for HBV markers (HBsAg, HBeAg, antiHBe, and HBV DNA) and for liver disease activity and severity (ALT, noninvasive tests of fibrosis-elastography, and/or liver biopsy) in order to identify patients for treatment and HCC surveillance. Serum HBV DNA level is important for the assessment of the phase of the infection, the decision to treat, and follow-up. The assessment of patients with chronic HBV infection is shown in Table 1. The first-degree relatives and sexual partners of individuals with chronic HBV infection should be tested for HBV serological markers (HBsAg, antiHBs, and antiHBc) and to be vaccinated, if negative [33].

Main treatment strategies of chronic HBV infection include 2 different concepts: therapy with nucleotide analog (NA) or with PEGylated-interferon-α (PEGInf α). The NA approved in Europe includes NA with low barrier against HBV resistance: lamivudine, adefovir, and telbivudine and NA with high barrier to HBV resistance: entecavir, tenofovir disoproxil fumarate, and tenofovir alafenamide.
The indication for treatment includes all patients with any detectable HBV DNA (2,000 IU/mL) and at least moderate liver inflammation, fibrosis, or cirrhosis (compensated or decompensated). Patients with HBV DNA 20,000 IU/mL and ALT ≥ 2x ULN should also be treated, as well as patients with HBV infection and family history of HCC, cirrhosis, or presence of extrahepatic manifestations of HBV infection [34].

The goal of treatment is the induction of long-term suppression of HBV DNA levels. The loss of HBeAg represents a partial immune control of the chronic HBeAg-positive individuals, while loss of HBsAg indicates profound suppression of HBV replication. A biochemical response to treatment is defined as ALT normalization and is usually achieved in patients with long-term suppression of HBV replication. Treatments with potent agents have beneficial effects in liver function, delay the occurrence of hepatic decompensation, and can improve outcomes and survival. HCC in patients with HBV infection is one of the main concerns and can develop in patients with an early stage of HBV infection or even in individuals who have been effectively treated [35].

**Hepatitis E Virus Infection**

Hepatitis E virus (HEV) infection is becoming a new European problem with at least 2 million locally acquired HEV in Europe, every year. HEV is mostly zoonotic infection with genotype 3 and genotype 4 that primarily spreads through pork meat products and causes silent epidemics in the “Western world.” Genotypes 1 and 2 spread through dirty water and mostly cause infection and often deaths in developing parts of the world [36]. Acute infection is usually clinically silent, with <5% of patients who develop symptoms such as fatigue, itching with elevated liver enzymes, and jaundice [37]. While immunocompetent patients with acute hepatitis E can clear the infection spontaneously, immunosuppressed patients may develop a chronic HEV infection. This has been seen only with HEV genotypes 3 and 4 [38].

HEV should be tested in all patients with hepatitis as part of the extended liver blood tests screening, patients presenting with DILI, unexplained flares of CLD, and all immunosuppressed patients with unexplained elevation of LFTs. Proposed diagnostic algorithm should consist of both serology and nucleic acid amplification techniques [39]. Positive laboratory tests that are commonly presented in acute HEV infection are as follows: HEV RNA with or without antiHEV IgM and/or antiHEV IgG or HEV antigen. Positive laboratory tests in chronic HEV infection can be as follows: HEV RNA ± antiHEV or HEV antigen, while past infection is determined by the presence of antiHEV IgG [36].

Acute HEV infection usually does not require therapy, except for the cases of severe acute hepatitis or acute-on-chronic liver failure. In such cases, ribavirin treatment may be considered. In transplant patients, ribavirin monotherapy is the treatment of choice together with reduction of the dose of immunosuppression, if possible. Patients who are nonresponders to ribavirin therapy should be offered PEGInf α for 3 months [36, 40].

**Autoimmune Hepatitis**

AIH is a relatively rare progressive, inflammatory CLD. AIH can affect any age, with peaks around puberty and between 40 and 60 years of age, and include both sexes and all ethnic groups. A third of patients are either asymptomatic or have severe acute fulminant hepatitis, while a third of patients already have cirrhosis at the time of diagnosis. Most common symptoms are fatigue, right upper quadrant pain, anorexia, itching, depression, jaundice, and polyarthralgia [41].

AIH is usually characterized by increased IgG levels, in association with human leukocyte antigens DR3 or DR4 and specific circulating autoantibodies. Depending on specific antibodies, it is subclassified into 3 major types: AIH type 1 (AIH-1) and AIH type 2 (AIH-2) and AIH type 3 (AIH-3). In AIH-1, antinuclear antibodies and/or smooth muscle autoantibodies are detected, and usually perinuclear anti-neutrophil cytoplasmic antibodies are also found. In AIH-2, specific autoantibodies, namely, anti-liver/kidney microsomal antibody type 1 or rarely anti-LKM type 3 and/or antibodies against liver cytosol type 1 antigen, can be detected. AIH-3 is very similar to AIH-1, with anti-SLA/LP antibodies, often Ro52-antibody is detected [42]. AIH-2 is more frequent in children and young adults, while AIH-1 is typically present in adults. AIH-2 may occur with acute and severe onset with advanced histological lesions at presentation. AIH-3 is similar to AIH-1 but has more severe clinical presentation [43]. Liver biopsy remains a gold standard for the diagnosis.

The first-line therapy is prednisolone in dose 0.5–1 mg/kg/day with azathioprine to a maintenance dose of 1–2 mg/kg/day [44]. Budesonide (9 mg/day) may be considered in noncirrhotic patients with early stage of disease.
An adequate therapeutic response to immunosuppression confirms the diagnosis.

HCC development in AIH is less common than other liver diseases. AIH patients with cirrhosis should undergo liver ultrasound in 6-month intervals for HCC screening [46].

**Drug-Induced Liver Injury**

DILI is classified as intrinsic (or direct) and idiosyncratic. Intrinsic DILI is dose related and occurs in a large proportion of individuals exposed to the drug (predictable), and onset is within a short time (hours to days). Acetaminophen hepatotoxicity is a prototype of intrinsic DILI and is the most common cause of acute liver failure in the USA and Europe [47]. Idiosyncratic DILI is usually not dose related, is unpredictable, and exhibits a variable latency to onset of days to weeks [48, 49]. Evidence from well-designed studies indicates that drugs work synergistically with other risk factors, age, sex, alcohol intake, T2DM, and obesity, contributing to pathogenesis and progression of liver disease [50–53]. Phenotypes of DILI and associated medications are shown in Table 2.

Liver biopsy is usually not required for diagnosis. Treatment involves withdrawal of the causative agent. Recovery is usually spontaneous without any specific treatment. The particular therapies are reserved for specific forms of DILI [54].

**Hereditary Hemochromatosis**

Hereditary hemochromatosis is one of the most common inheritable genetic disorders. One in 200 individuals in northern Europe is affected by this condition. Polymorphisms in the HFE gene (most frequent homozygotes for the C282Y polymorphism) cause iron overload due to excess iron absorption and induce accumulation in tissues and organs. Without appropriate treatment, excess iron leads to cirrhosis while one-third of patients will develop a primary liver cancer [55].

Table 2. Phenotypes of DILI and associated medications

<table>
<thead>
<tr>
<th>Phenotypes of DILI</th>
<th>Medications</th>
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<tbody>
<tr>
<td>Idiosyncratic DILI</td>
<td>Antibiotics: amoxicillin-clavulanate, erythromycin, flucloxacillin, interferon alpha/peginterferon, isoniazid, ketoconazole, minocycline, nevirapine, nitrofurantoin, pyrazinamide, rifampicin, co-trimoxazole, and sulfonamides</td>
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<td></td>
<td>Central nervous system: carbamazepine, chlorpromazine, dantrolene, halothane, phenytoin, and valproate</td>
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<td></td>
<td>Cardiovascular: amiodarone, hydralazine, methyldopa, quinidine, statins (atorvastatin and simvastatin)</td>
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<td></td>
<td>Immunomodulatory: azathioprine/6-mercaptopurine, interferon beta, methotrexate, and thioguanine</td>
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<td></td>
<td>Antineoplastic: busulfan, flusuridine, and flutamide</td>
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<td></td>
<td>Rheumatologic: allopurinol, auranofin/gold products, diclofenac, ibuprofen, nimesulide, and sulindac</td>
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<td></td>
<td>Endocrine: anabolic androgenic steroids, estrogens/progestins, and propylthiouracil</td>
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<td></td>
<td>Others: disulfiram and ticlopidine</td>
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<tr>
<td>Drug-induced autoimmune hepatitis</td>
<td>Diclofenac, halothane, indomethacin, infliximab, methyldopa, minocycline, nitrofurantoin, and statins</td>
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<tr>
<td>Secondary sclerosing cholangitis</td>
<td>Amiodarone, atorvastatin, amoxicillin-clavulanate, gabapentin, infliximab, 6-mercaptopurine, sevoflurane, and venlafaxine</td>
</tr>
<tr>
<td>Granulomatous hepatitis</td>
<td>Allopurinol, carbamazepine, methyldopa, phenytoin, quinidine, and sulphonamides</td>
</tr>
<tr>
<td>Acute fatty liver</td>
<td>Amiodarone, didanosine, stavudine, valproate, zalcitabine</td>
</tr>
<tr>
<td>Drug-associated fatty liver disease</td>
<td>Methotrexate, 5-fluorouracil, irinotecan, tamoxifen, corticosteroids, lomiptamide, and mirpanser</td>
</tr>
<tr>
<td>Nodular regenerative hyperplasia</td>
<td>Azathioprine, busulphan, bleomycin, cyclophosphamide, chlorambucil, cysteine arabinoside, carmustine, doxorubicin, 6-thioguanine, and oxaliplatin</td>
</tr>
<tr>
<td>Ductopenic (vanishing bile duct) syndrome</td>
<td>Azathioprine, androgens, amoxicillin-clavulanate, carbamazepine, chlorpromazine, erythromycin, estradiol, flucloxacillin, phenytoin, terbinafine, and co-trimoxazole</td>
</tr>
<tr>
<td>Liver tumors</td>
<td>Anabolic androgenic steroids and oral contraceptives</td>
</tr>
</tbody>
</table>

DILI, drug-induced liver injury.
Wilson’s Disease

Wilson’s disease in another inherited condition in which defective biliary excretion of copper leads to copper accumulation in the liver and brain. It is a rare autosomal recessive disorder with a frequency of 1 in 30,000. About 3% of patients present beyond the fourth decade, either with hepatic or neurologic disease. Clinical presentation can vary widely, but most of the patients develop cirrhosis, neuropsychiatric disturbances, and Kayser-Fleischer rings, while some present with the acute liver failure and acute onset of hemolysis [56].

Hepatocellular Carcinoma

HCC represents about 90% of primary liver cancers and constitutes a major global health problem. HCC is the sixth most common cancer in the world and the third most frequent cause of cancer-related death globally, while incidence is rapidly rising in last decades [57].

Major risk factors include infections HCV and HBV, alcohol, and aflatoxin exposure. Only 40% of patients with HCC are diagnosed at early stages. Patients with cirrhosis are at the highest risk and should undergo liver US for HCC screening in every 6 months. Serological tests including alpha fetoprotein are not currently cost effective. Diagnosis of HCC in cirrhotic patients should be based on noninvasive criteria and/or pathology. Surgical resection is the treatment of choice in patients with HCC arising on a noncirrhotic liver [58].

There are several different algorithms in the treatment of HCC. One of the most widely used is the Barcelona Clinic Liver Cancer (BCLC) system, which classifies patients into 5 stages. For very early-stage HCC or stage 0 single tumor, <2 cm (carcinoma in situ), resection is indicated. For very early-stage HCC or stage 0 and for early stage or stage A HCC (no >3 nodules, each <3 cm), in the absence of liver disease, LT is the therapy of choice. For stage 0 and stage A in the presence of associated liver diseases, radiofrequency ablation is the therapy of choice. For intermediate or stage B HCC (multinodular), transarterial chemoembolization is recommended. For advanced or stage C HCC (portal invasion, N1, and M1), sorafenib is recommended. For terminal or stage D disease (Child-Pugh C), palliative treatment with supportive care is recommended [59].

Diagnosing benign liver tumors and differentiating in relation to HCC is based on radiological examination, and it is primarily related to the size of the lesion. Lesions <1 cm are monitored by ultrasound at 4 months. For lesions >1 cm, CT or MR examination should be performed [60].

Conclusion

CLD is amenable to prevention and treatment, reducing the burden of liver disease and deaths due to end-stage liver diseases. Removal of the etiological factors causing liver injury is the most important step in the management of cirrhosis. To prevent complications, patients with CLD need surveillance. Treatment with potent agents for some CLD has beneficial effects in liver function, delays the occurrence of hepatic decompensation, and improves outcome and survival. The integrated approach is needed to achieve the best possible outcomes including the primary care physician, gastroenterologist/hepatologist, and liver transplant team to closely monitor patients with CLD in order to prevent complication and timely recognition for LT.

Conflict of Interest Statement

The authors have no conflicts of interest to declare.

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Author Contributions

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