Review Article

Hepatitis B Virus: Update in Management

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Summary

Hepatitis B virus (HBV) infection is a major public health problem worldwide. HBV, a member of the hepadnaviridae family, is a small DNA replicates through an RNA intermediate and can integrate into the host genome. It is transmitted through contact with infected blood and semen. HBV infection leads to a wide spectrum of liver disease ranging from acute - fulminant hepatic failure to chronic hepatitis, cirrhosis, and hepatocellular carcinoma. Infants and children are more likely to develop a chronic hepatitis B infection, while most adults infected with the virus recover even if their signs and symptoms are severe, but 5%-10% are unable to clear the virus and become chronically infected. Screening for HBV infection could identify chronically infected persons who may benefit from treatment or other interventions, such as surveillance for hepatocellular carcinoma. Most people diagnosed with chronic hepatitis B infection need treatment for the rest of their lives. The primary treatment goals for patients with HBV infection are to prevent progression of the disease to cirrhosis, liver failure, and hepatocellular carcinoma. Moreover, prevents infection from passing to others. Treatment for chronic hepatitis B may include several antiviral medications such as, entecavir, tenofovir, lamivudine, adefovir and telbivudine. These drugs can help fight the virus and slow its ability to damage your liver. WHO recommends that all infants receive the hepatitis B vaccine as soon as possible after birth, preferably within 24 hours?

Keywords: Hepatitis B virus, Nucleoside/nucleotide analogues and hepatitis B vaccine

Introduction

Infection with hepatitis B virus (HBV) remains an important global public health problem with significant morbidity and mortality. The prevalence of chronic HBV infection is highest in the Western Pacific (6.2%) and African regions (6.1%). The Eastern Mediterranean region has high prevalence (3.3%); the prevalence is lower in the South–East Asian (2.0%) and European regions (1.6%) and is lowest in the North and South American regions (0.7%). The prevalence is decreasing in several endemic countries due to improvements in the socioeconomic status, universal vaccination programs and perhaps effective antiviral treatments. There are ten different genotypes of HBV (A-J), and the geographical distribution of each HBV genotype is distinct. Infection with different genotypes of HBV is associated with different outcomes in chronicity, disease progression and responses to IFNα treatment; however, the approved HBV vaccines are effective against all genotypes.

Natural history of HBV infection

New terminology has been suggested for the phases of chronic HBV infection. The previously used terms of immune tolerance, immune active and/or clearance and immune control and/or residual phases have been changed. HBV infection can be classified now into five phases, Tab. (1): (I) HBeAg-positive chronic infection, (II) HBeAg-positive chronic hepatitis, (III) HBeAg-negative chronic infection, (IV) HBeAg-negative chronic hepatitis and (V) HBsAg-negative phase.
**Phase 1: HBeAg-positive chronic HBV infection.**

Previously named immune tolerance phase. This phase (also termed the high replicative, low inflammatory phase) typically occurs in patient infected perinatally. It is characterized by high viral load, near normal liver histology owing to a minimal host immune mediated reaction. ALT persistently within the normal range according to traditional cut-off values [upper limit of normal approximately 40 IU/L] and serum positivity for HBeAg. Recent studies have indicated that adolescents may have detectable HBV-specific functionally active T cells. Studies of this phase of the disease have also revealed clonal expansion of hepatocytes and visible HBV integrands, which may contribute to the development of HCC by transactivating human oncoproteins. These patients are highly infectious owing to the high levels of HBV DNA.

**Phase 2: HBeAg-positive chronic hepatitis B.**

The second phase, historically termed immune clearance or immune active phase, most commonly occurs in the third or fourth decade of life in patients who were infected with HBV at an early age. The prominent features of this phase are the presence of serum HBeAg, high levels of HBV DNA and elevated ALT and presence of moderate-severe liver necroinflammation. In this phase, the triggering of immune T cell-mediated responses against infected hepatocytes leads to the cytolysis release of alanine transaminases (ALT) and reduction of HBV DNA levels. However, the intensity of the immune response fluctuates over time, resulting in fluctuating levels of ALT (hepatitis flares, which are defined differently in different studies; for example, twofold elevation from the baseline ALT levels and HBV DNA. This phase can be of varying duration and ends with a reduction of HBV DNA level and HBeAg seroconversion to anti-HBe positivity and enter the HBeAg-negative infection phase in majority of patients. Other patients may fail to control HBV and progress to the HBeAg-negative CHB phase for many years. A proportion of patients (annual rate of ~15%) will spontaneously seroconvert from HBeAg to anti-HBe; these patients (previously referred to as inactive carriers) typically show absence of serum HBeAg with detectable anti-HBe, near normal serum aminotransferases and HBV DNA concentrations of <2,000 IU per ml. Certain patients in this phase, still have HBV DNA levels >2,000 IU/ml however <20,000 IU/ml associated with persistently normal ALT and only minimal hepatic necroinflammatory activity and low fibrosis. These patients have low risk of progression to cirrhosis or HCC if they remain in this phase. In a substantial proportion of patients in Asian and Mediterranean countries, HBeAg seroconversion is associated with the occurrence of precore and/or core promoter mutations. In these patients, HBeAg is absent because of the selection of HBV virions that do not express HBeAg (precore mutant HBV). Finally, patients may have spontaneous HBsAg sero clearance during the natural course of disease. Various studies have shown that patients with spontaneous HBsAg sero clearance at an earlier age have a better outcome than those without HBsAg sero clearance.

**Phase 4: Named HBeAg-negative chronic hepatitis B.**

It is characterised by the absence of serum HBsAg usually with detectable anti-HBe, and persistent or fluctuating moderate to high levels of serum HBV DNA (often lower than in HBeAg-positive patients), in addition to fluctuating or persistently elevated ALT values. The liver histology shows necroinflammation and fibrosis. The majority of these patients harbour HBV variants in the precore and/or the basal core promoter regions that abate or stop HBeAg expression. This phase is associated with low rates of spontaneous disease remission.

**Phase 5: HBsAg-negative phase**

It is characterised by serum negative HBsAg and positive antibodies to HBeAg (anti-HBe), with or without detectable antibodies to HBsAg (anti-HBs). This phase is also known as “occult HBV infection”. In rare cases, the absence of HBsAg could be related to the sensitivity of the assay used for detection. Patients in this phase have normal ALT values and usually, but not always, undetectable serum HBV DNA. HBV DNA (cccDNA) can be detected frequently in the liver. HBsAg loss before the onset of cirrhosis is associated with a minimal risk of cirrhosis, decompensation and HCC, and an improvement on survival. However, if cirrhosis has developed before HBsAg loss, patients
remain at risk of HCC therefore, HCC surveillance should continue. Immunosuppression may lead to HBV reactivation in these patients. Table (1) Natural history of patients with chronic HBV infection.

<table>
<thead>
<tr>
<th>HBsAg status</th>
<th>Chronic infection</th>
<th>Chronic hepatitis</th>
<th>Chronic liver disease</th>
</tr>
</thead>
<tbody>
<tr>
<td>HBsAg+</td>
<td>High</td>
<td>High</td>
<td>Low</td>
</tr>
<tr>
<td>HBsAg-</td>
<td>Negative</td>
<td>Negative</td>
<td>Negative</td>
</tr>
<tr>
<td>HBV DNA</td>
<td>&gt;10^5 IU/ml</td>
<td>&lt;10^5 IU/ml</td>
<td>&lt;200 IU/ml</td>
</tr>
<tr>
<td>ALT</td>
<td>Normal</td>
<td>Elevated</td>
<td>Elevated</td>
</tr>
<tr>
<td>Liver disease</td>
<td>Normal</td>
<td>Moderately severe</td>
<td>Severe</td>
</tr>
<tr>
<td>Age in years</td>
<td>Immature</td>
<td>Mature</td>
<td>Old</td>
</tr>
</tbody>
</table>

*Persistently or intermittently. **HBV DNA levels can be between 2,000 and 20,000 IU/ml in some patients without signs of chronic hepatitis.

Diagnosis, Screening and Prevention

Diagnosis

The key serological marker of acute and chronic hepatitis B is the detection of HBsAg in serum. However, several other HBV serological markers are clinically useful in HBV infection. For example, in patients with clinical recovery of HBV infection, anti-HBs and hepatitis B core antigen (HBcAg; anti-HBc) may be detectable, tab. (2). In addition to serum tests, an assessment of the fibrosis and cirrhosis status in patients with chronic HBV infection is also important for disease progression, treatment indication and management. An accurate but invasive test of liver disease is liver biopsy, whereas noninvasive tests include transient elastography (to measure liver stiffness) or various serum biomarkers. With the availability of the latter two methodologies, most patients and clinicians are reluctant to go through with liver biopsy because of its invasive nature. Algorithms for the use of serum biomarkers and liver stiffness measurements (if transient elastography is available) can be used in clinical practice to adjudge the degree of fibrosis in chronic hepatitis B.

Detection of HBeAg

HBeAg serological testing, usually performed in chronic HBV-infected patients, distinguishes two forms of hepatitis B: HBeAg-positive and HBeAg-negative chronic HBV infection. HBeAg seroconversion marks the transition from the immune clearance phase or HBeAg-positive chronic hepatitis to the immune control phase or HBeAg-negative chronic infection or hepatitis.

Detection of anti-HBc

Total anti-HBc measured by immunoassay is a marker of acute, chronic and resolved HBV infection, or occult hepatitis B. In addition, the presence of anti-HBc can predict reactivation of HBV associated with rituximab or immunosuppressive therapy. Immunoglobulin M (IgM) anti-HBc is detected during acute HBV infection and is also detected during exacerbations of chronic HBV, and importantly, it may be the only serological marker detectable in fulminant acute hepatitis B. Thus, measurement of anti-HBc is usually recommended for diagnosis in patients suspected to have an acute exacerbation of chronic HBV infection or to decide if patients who are planning to undergo immunosuppressive therapy will require prophylactic antiviral therapy.

Detection of HBV DNA

HBV DNA tests should be performed on a regular basis (every 6 months) in all patients with chronic HBV. The concentrations of HBV DNA provide an indication for therapy within current guidelines, and for monitoring treatment efficacy. Indications for treatment are based on the association between HBV DNA concentrations and the risk of HCC. A decline in HBV DNA concentrations can predict the efficacy of treatment, and an increase in HBV DNA in serum is observed with the development of resistance to NUCs.
Co-morbidities
It including other causes of chronic liver disease should be systematically excluded including co-infections with hepatitis D virus (HDV), hepatitis C virus (HCV) and HIV.

Testing for antibodies
Testing for antibodies against hepatitis A virus should be performed, and patients with negative anti-HAV should be advised to be vaccinated against HAV.

Table (2) Interpretation of screening tests for HBV infection

<table>
<thead>
<tr>
<th>HBsAg</th>
<th>IgM</th>
<th>Anti-HBc</th>
<th>Anti-HBs</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>+</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>Acute infection</td>
</tr>
<tr>
<td>+</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>Chronic infection</td>
</tr>
<tr>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>Chronic infection</td>
</tr>
</tbody>
</table>

HBsAg= hepatitis B surface antigen; anti-HBc = antibody to hepatitis B core antigen; anti-HBs = antibody to HBsAg; IgM = immunoglobulin M; mIU/mL = milli-International Units per milliliter.

HBV Vaccine Indications
The WHO recommends universal vaccination for HBV and that all infants should receive the first dose of vaccine soon after birth, preferably within 24 hours. The birth dose, or vaccination within the expanded programme on immunization (EPI) dose, is followed by an additional two to three doses, depending upon national policies and prevalence, tab. (3).

Table (3) High-risk unvaccinated individuals for which hepatitis B vaccine is recommended

Goals of therapy
The ultimate goal is total eradication of HBV infection by various strategies, including vaccination, treatment and prevention of transmission. The goal of therapy for chronic HBV infection is to improve quality of life and survival of the infected person via preventing progression of the disease to cirrhosis, decompensated cirrhosis, end-stage liver disease, HCC and death. Also to prevent transmission of HBV to others, hepatitis B reactivation and the prevention and treatment of HBV-associated extrahepatic manifestations.

This goal can be achieved by suppression of HBV replication in sustained manner. Regression of fibrosis and cirrhosis can be regarded as a further goal of treatment in patients with established advanced fibrosis or cirrhosis. In patients with HBV-induced HCC, the goals of nucleos(t)ide analogue (NA) therapy are primarily to suppress HBV replication to induce the stabilization of HBV-induced liver disease and to prevent disease progression, and to reduce the risk of HCC recurrence after potentially curative HCC therapies. Stabilizing the HBV-induced liver disease can also be regarded as a prerequisite for the safe and effective applications of HCC treatments.

End points of treatment
HBV treatment is typically aimed to achieve profound virological suppression, which in turn will lead to biochemical remission (return of ALT values to the normal range), histological improvement and prevention of the complications of liver disease. The ideal endpoint in both HBeAg-positive and HBeAg-negative patients is sustained off-therapy HBsAg loss, with or without seroconversion to anti-HBs. This is associated with a complete and definitive remission of the activity of CHB and an improved long-term outcome. This endpoint, however, is infrequently achievable with the currently available anti-HBV agents. A more realistic endpoint is the induction of sustained or maintained virological remission. Induction of sustained off-therapy virological response in both HBeAg-positive (with sustained anti-HBe seroconversion) and HBeAg-negative patients is a satisfactory endpoint, because it has been shown to be associated with improved prognosis. If sustained off-therapy response not ach-
iable, then a maintained virological remission (undetectable HBVDNA by a sensitive PCR assay) under long-term antiviral therapy in HBeAg-positive patients who do not achieve anti-HBe seroconversion and in HBeAg-negative patients is the next most desirable endpoint. A more advanced functional cure is HBsAg seroclearance, as spontaneous or treatment-associated HBsAg seroclearance substantially decreases the risk of HCC development, provided the patient is <50 years of age, and has not developed cirrhosis. However, HBsAg seroclearance is rarely achieved with the currently available anti-HBV agent’s. Current potent NUCs have a profound effect on virological suppression with continued therapy. For NUCs: a virological response is defined as undetectable serum HBV DNA (10–60 IU per ml); a serological response as HBeAg and/or HBsAg seroconversion; a biochemical response as normalization of serum ALT level; and a histological response as a decrease in necroinflammation of the liver. Worldwide, Pegylated interferon-α (PEG-IFNα) therapy is given for a finite duration to a small subset of patients (discussed below) with the aim of achieving sustained off-treatment immune control. The response is assessed 6 months post-treatment, with criteria similar to those of NUCs, except for the virological response, which is defined as HBV DNA <2,000 IU per ml. Long-term follow-up studies of sustained treatment with NUCs demonstrate that a profound reduction in development of cirrhosis can be achieved with the early initiation of treatment. Consequently, this has resulted in an almost complete elimination of liver failure in the setting of treated patients with chronic HBV infection. In addition, there is a clear reduction, but not complete prevention, of the development of HCC.

**Treatment Agents**

Treatment agents approved by the US FDA, European Medicines Agency and many countries in Asia are broadly classified into immunomodulatory agents and antiviral agents, tab. (4). The former includes conventional IFNα-2b and PEG-IFNα-2a. The latter includes (nucleoside/nucleotide analogues) NUCs, namely:

*) Lamivudine (LAM)
*) Adefovir dipivoxil (ADV)
*) Entecavir (ETV)
*) Telbivudine (TDF)
*) Tenofovir alafenamide (TAF).

The mainstay of treatment in most countries is NUCs. Of the six NUCs licensed by FDA for HBV treatment, ETV, TDF and TAF are the advocated first-line agents because of their potency and low resistance rates. Entecavir has negligible adverse effects and is an excellent agent for treatment-naive patients. However, patients previously treated with lamivudine who have developed lamivudine resistance will already have two of the three mutations required for the development of entecavir resistance; 51% of these individuals will develop entecavir resistance within 5 years. In lamivudine-resistant patients, TDF and TAF can achieve a higher intrahepatic level of tenofovir; thus, a lower dose of TAF can be used to exert the same degree of viral suppression. As such, the off-target effects of TAF are decreased.

According to one study, TAF has comparable viral suppression to TDF with improved rates of ALT normalization, substantially fewer declines in hip and spine bone mineral density (side effect of TDF) and smaller decreases in estimated glomerular filtration rate, indicating TAF has less of an off-target effect on renal function. However, in general, clinical adverse effects are considered to be minimal during long-term NUC therapy.

**Table (4) Approved HBV antiviral therapy in children and adults**

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dose in Adults</th>
<th>Use in Children</th>
<th>Pregnancy Category</th>
<th>Potential Side Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tenofovir</td>
<td>300 mg daily</td>
<td>&gt;12 years</td>
<td>C</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td>15 mg daily</td>
<td>2–12 years</td>
<td>B</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td>5 mg daily</td>
<td>—</td>
<td>B</td>
<td>N/A</td>
</tr>
<tr>
<td>Nucleoside analogues</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lamivudine</td>
<td>100 mg daily</td>
<td>&gt;12 years</td>
<td>C</td>
<td>N/A</td>
</tr>
<tr>
<td>Adefovir</td>
<td>10 mg daily</td>
<td>&gt;12 years</td>
<td>C</td>
<td>N/A</td>
</tr>
<tr>
<td>Telbivudine</td>
<td>600 mg daily</td>
<td>&gt;12 years</td>
<td>C</td>
<td>N/A</td>
</tr>
</tbody>
</table>

*Dose adjustments are needed in patients with renal dysfunction.* In 2015, the U.S. Food and Drug Administration replaced the pregnancy risk designation by letters A, B, C, D, and X with more specific language on pregnancy and lactation. This new labeling is being phased in gradually.
and to date only TAF includes these additional data. Peg-IFN-a-2a is not approved for children with chronic hepatitis B, but is approved for treatment of chronic hepatitis C. Providers may consider using this drug for children with chronic HBV. The duration of treatment indicated in adults is 48 weeks. Entecavir dose is 1 mg daily if the patient is Lamivudine experienced or if they have decompensated cirrhosis. Abbreviation: TSH, thyroid stimulating hormone.

**Indications of treatment**

The indications for treatment are based largely on the combination of three criteria:
- Serum HBV DNA levels
- Serum ALT levels
- Severity of liver disease (assessed by clinical evaluation, liver biopsy or noninvasive methods).

Indications for treatment should also take into account age, health status, family history of HCC or cirrhosis and extrahepatic manifestations, tab. (5)\(^5,18,25\).

**Table (5) Indications for initiating treatment in chronic HBV infection.**

<table>
<thead>
<tr>
<th>Guideline</th>
<th>Individual who is HBeAg positive</th>
<th>Individual who is HBeAg negative</th>
</tr>
</thead>
<tbody>
<tr>
<td>AASLD 2017</td>
<td>HBV DNA IU/ml</td>
<td>ALT UI</td>
</tr>
<tr>
<td>EASL 2017</td>
<td>≥2,000</td>
<td>&gt;ULN or at least moderate liver necro-inflammatory or fibrosis</td>
</tr>
<tr>
<td>AASLD 2018</td>
<td>&gt;10,000</td>
<td>&gt;2xULN or significant histological disease</td>
</tr>
<tr>
<td>APASL 2017</td>
<td>≥30,000</td>
<td>&gt;2xULN or significant histological disease</td>
</tr>
</tbody>
</table>

**AASLD,** American Association for the Study of Liver Diseases; **ALT,** alanine aminotransferase; **APASL,** Asian Pacific Association for the Study of Liver; **EASL,** European Association for the Study of the Liver; **HBeAg,** hepatitis B e antigen; **HBV,** hepatitis B virus; **IU,** international units; **ULN,** upper limit of normal. A patient with cirrhosis with detectable HBV DNA should be treated irrespective of the HBV DNA and ALT levels.

Although, the liver biopsy is still to be the gold standard to estimate disease activity in CHB\(^26\), the treatment decisions may be based on non-invasive markers of fibrosis. Elastography is seemed to have the highest diagnostic accuracy for the detection of advanced fibrosis. The results of elastography should be interpreted with caution in patients with severe inflammation and high ALT level\(^27\). Patients with HBV DNA levels >20 000 IU/mL and ALT >2xULN can begin treatment without a liver biopsy because usually the results will not change the decision for treatment\(^25\). A non-invasive technique is recommended in these cases to confirm or exclude cirrhosis in patients who begin treatment without a liver biopsy\(^25\). Treatment may be begun in patients with HBV DNA levels >2000 IU/mL and at least moderate fibrosis, even if ALT levels are normal. In addition, patients with HBeAg-positive chronic HBV infection (persistently normal ALT and high HBV DNA levels) may be treated if they are >30 years old whatever the severity of histological lesions in the liver\(^25\). Since the course of chronic HBV infection fluctuates, patients who do not fulfill the indications for antiviral therapy should be carefully monitored with periodic assessment of serum ALT and HBV DNA levels as well as for the severity of liver fibrosis by non-invasive markers. Patients with HBeAg-positive chronic HBV infection who are <30 years old and not being treated should undergo ALT tests every 3 months at least, HBV DNA every 6-12 months and assessment of liver fibrosis every 12 months\(^25\). Patients with HBeAg-negative chronic HBV infection and HBV DNA <2000 IU/mL should be monitored with ALT measurements every 12 months and have HBV DNA and liver fibrosis tests every 2-3 years\(^28\). Recent data suggest that, quantitative HBsAg testing can be helpful in deciding on the frequency of follow-up in such patients. Patients with HBsAg levels <1000 IU/mL should undergo ALT measurements every 12 months and have HBV DNA and liver fibrosis tests every 3 years, while those with HBsAg levels ≥1000 IU/mL should receive ALT measurements every 6 months and HBV DNA and liver fibrosis tests every 2 years at least\(^25\). Patients with HBeAg-negative chronic HBV infection, normal ALT and HBV DNA ≥2000 IU/mL should be closely followed with ALT measurements every 3 months for the first year and every 6 months at least thereafter. Moreover, annual evaluation of HBV DNA levels and a non-invasive liver fibrosis test should be performed for at least 3 years. If they do not fulfill any indications for treatment during the first 3 years of follow-up, these patients should be monitored for life, like all other patients in this phase of disease\(^26\). Patients with decompensated cirrhosis and detectable HBV DNA, tab. (6) require urgent antiviral treatment with NA(s). Significant clinical improvement can be associated with control of viral replication\(^29,30\). However, the
patients should be considered for liver transplantation at the same time if antiviral treatment not sufficient to correct the decompensating state.\(^{18}\) Patients with compensated cirrhosis and HBV DNA $> 2000$ U/ml should also be considered for treatment even if ALT levels are normal.\(^{18}\)

Table (6) Treatment indications for chronic HBV-infected patients with cirrhosis or reactivation of chronic HBV infection.\(^{18}\)

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**References**


17- Centers for Disease Control and Prevention (CDC). Use of hepatitis B vaccination for adults with diabetes mellitus:


