Treatment options for hepatitis B

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Hepatitis B virus (HBV) is an enveloped DNA virus with a double-stranded circular genome that replicates through an RNA intermediate within the host hepatocyte (1-3). The virus infects the hepatocyte and is converted to covalently closed circular DNA, which serves as a template for viral replication. When the replication cycle of HBV is complete, mature, infectious virions are released into the bloodstream (1-3). The virus itself is noncytopathic, and the mechanism of cellular injury is immune-mediated. Viral clearance is mediated by both cytopathic and noncytopathic pathways and, although the majority of individuals are able to clear the virus and exhibit firm immunological control, some individuals fail to mount an adequate immune response to eliminate the virus, which leads to chronic infection (3,4). Chronic infection appears to be associated with inadequate or exhausted T cell responses to the virus, such that most patients with chronic HBV infection have few or no HBV-specific T cells in their circulation (2). The use of nucleoside analogue treatments may transiently restore T cell responses (5,6).

Hepatitis B remains a major health concern worldwide. It is the leading cause of chronic hepatitis, cirrhosis and hepatocellular carcinoma (HCC). It is estimated that 400 million people worldwide are infected with HBV, and in 2005, the World Health Organization reported that during the 2002 calendar year, over 600,000 patients worldwide died from HBV or its complications (7). According to estimates from the United States, 100,000 people per year become infected, and up to 5000 people per year die as a result of HBV or its complications (8). The incidence rate of clinically recognized acute hepatitis B in Canada has been estimated to be 2.3 per 100,000 people, which equates to approximately 700 cases per year, although the figure may be as high as 3000 cases per year (9,10). In Canada, the exact prevalence of chronic hepatitis B is generally unknown, but HBV infection has been estimated to affect 0.5% to 1% of the population, dependent largely on the prevalence of HBV within the immigrant populations (10). Developments in nucleoside analogue therapy that have occurred over the past decade, coupled with our increasing understanding of the natural history of HBV infection, provided an impetus for an overview of new developments in this area.

Approximately one in 20 adults and nine in 10 infants acutely infected with HBV will develop chronic infection (1,2). Of those with acute infection, approximately one in three adults have symptomatic hepatitis. The major determinant of outcome appears to be immunological failure to clear the virus, typified by persistence of detectable HBV DNA levels and hepatitis B e antigen (HBeAg) in serum (4). Additionally, viral heterogeneity may also play a role in the outcome. Eight HBV genotypes have been described, with genotype A being most common in northern Europe and North America, genotypes B and C more commonly seen in Asia, and genotype D being more common in the Mediterranean (1,2). Individuals infected with genotype C appear to have a greater likelihood of progression to cirrhosis and the development of HCC. Chronic HBV infection may be divided into phases based on alanine aminotransferase (ALT) levels, presence of HBeAg, HBV DNA levels and immune status (2). These phases are referred to as the inactive carrier state, the immune-tolerant phase, HBeAg-positive (HBeAg+) chronic hepatitis phase and HBeAg-negative (HBeAg–) chronic hepatitis phase (2). The inactive carrier state (ie, low or nondetectable HBV DNA levels, absence of HBeAg, presence of hepatitis B surface antigen [HBsAg], normal ALT levels and nonactive liver histology) usually has a benign course but can reactivate or transition into the HBeAg– chronic hepatitis phase (11). Although the course of the inactive carrier state is typically benign, depending on the time taken to achieve this phase, there are various amounts of preceding hepatic inflammation that can lead to fibrosis, including the potential for cirrhosis. The inactive carrier state does not necessarily preclude the presence of liver disease and the development of cirrhosis and HCC. The HBeAg– chronic hepatitis phase, typified by moderately elevated HBV DNA levels, absence of HBeAg, presence of HBsAg, elevated ALT levels and active liver histology, leads to progressive liver disease with its attendant complications. The HBeAg+ chronic hepatitis phase is the usual pattern and is typified by highly elevated HBV DNA levels, presence of HBeAg and HBsAg, elevated ALT levels and active liver histology, and leads to progressive liver disease with its attendant complications. A proportion of individuals in the HBeAg+ chronic hepatitis phase have no elevations of ALT levels or nonactive liver histology despite high levels of HBV DNA, and are referred to as being in the immune-tolerant phase (12). This phase of the disease has not been associated with progressive liver disease.

The natural history of chronic HBV infection is also impacted by a number of factors other than the phase of the disease, and may be quite variable depending on a complex interplay between viral replication and host immune response in any given individual. Disease progression may be impacted by genotypes or mutations, mode of transmission, age, alcohol consumption, obesity and concurrent viral infections such as HIV or hepatitis C virus. Large, long-term natural history studies of HBsAg+ persons have been conducted in Asia and Europe. These studies, representing the best data to date, have shown that the development of cirrhosis and HCC is
frequent and correlates with several pre-existing host and viral factors. The largest and most thorough evaluation was in a population-based survey by Chen et al. (13), in which HCC developed in 4.3% of 3653 HBsAg+ persons (0.4% per year) who were followed for an average of 11.4 years. Risk factors for HCC included male sex (hazard ratio [HR] = 3.0), older age (HR = 3.6 to 8.3), cigarette smoking (HR = 1.7) and alcohol consumption (HR = 2.6). Viral and disease factors predictive of HCC development included elevations in serum ALT levels (HR = 4.1), presence of HBeAg (HR = 4.2) and higher levels of HBV DNA. Levels of HBV DNA at the time of initial evaluation were most closely linked with the eventual development of HCC and in a dose-dependent manner, levels above 100,000 copies/mL were strongly linked (HR = 8.9 to 10.7) and levels above 10,000 copies/mL were significantly linked (HR = 2.7). The relationship between HBV DNA levels and HCC held true, even for patients with normal ALT levels at the time of initial evaluation. This study has been criticized for its generalization to non-Chinese populations that are not male and elderly, although almost 40% of study patients were female, 90% were 59 years of age or younger, and 71% were 49 years of age or younger.

A patient may be considered to have chronic HBV if HBsAg is still detected after six months, hepatitis B surface antibody is not detectable, serum HBV DNA level is elevated, aspartate aminotransferase (AST) and ALT levels are persistently or intermittently elevated and liver biopsy demonstrates chronic hepatitis. However, a normal ALT level does not exclude significant liver disease. The development of chronic infection has no predictable timeframe and can occur immediately after the acute phase of HBV infection or several years later. Unfortunately, the infection may remain silent for many years before symptoms and signs of cirrhosis become evident. Therefore, the major goals of therapy are long-term prevention of progression to cirrhosis, end-stage complications of cirrhosis and HCC (2, 14, 15). The decision to provide antiviral therapy for individuals with chronic hepatitis B has evolved over time. Current therapy recommendations (2, 14-16) are based on several factors—age, ethnicity, HBV DNA levels, HBeAg status, ALT levels, histological picture and comorbidity. For those patients with fibrosis and cirrhosis, antiviral therapy is indicated at any detectable HBV DNA level. HBeAg+ patients with HBV greater than 20,000 U/mL and HBeAg– patients with HBV greater than 2000 U/mL with increased ALT levels and patients with normal ALT levels but active necroinflammation on liver biopsy are candidates for therapy. Markers used to assess response to treatment include normalization of ALT levels, loss of HBV DNA and improved liver histological findings (14-16). Complete eradication of HBV is difficult because the virus tends to integrate into the host genome. Therapies can be categorized as either immunomodulators or antivirals.

The two agents available to modulate the immune response in patients with HBV infection are interferon alpha-2b (IFN-α2b), and pegylated IFN-α2a (PEG IFN-α2a) and PEG IFN-α2b. Currently, IFN-α2b and PEG IFN-α2a are indicated for treatment in HBeAg+ and HBeAg– patients with chronic hepatitis B and are administered subcutaneously once daily, three times a week or once weekly, respectively. The recommended treatment duration for HBeAg+ patients is 16 to 24 weeks (14-16). Approximately 25% to 40% of HBeAg+ patients treated with IFN-α2b experience a loss of serum HBV DNA or HBeAg after 12 to 24 weeks of therapy. Predictors of poor response include a high HBV DNA level, age older than 40 years, male sex and cirrhosis (14-16). PEG IFN-α2a, with a longer half-life, lacks the peaks and troughs associated with traditional interferon and is more convenient to administer. Chronic HBV guidelines (14-16) recommend at least 12 months of treatment for HBV+ patients. A major drawback of treating HBeAg– patients with IFN-α2b is that responses are less durable than those of HBeAg+ patients, with only 19% of patients having undetectable HBV DNA levels 24 weeks after stopping therapy (14-16). Interferon therapy, both the traditional and pegylated versions, are associated with a wide array of adverse events, including flu-like symptoms, which may occur in up to 90% of patients. Although patients may develop tolerance to some of these side effects after the first week of therapy, fatigue, anorexia, alopecia, mood swings, anxiety and depression may occur throughout treatment.

Antiviral agents may be classified as L-nucleosides (including lamivudine, emtricitabine and telbivudine), acyclic phosphonates (including adefovir and tenofovir) or cyclopentenones (entecavir) (2). Emtricitabine and tenofovir are not yet approved for use for chronic HBV infections in Canada.

Lamivudine, a nucleoside analogue, was the first antiviral agent approved for treatment of chronic hepatitis B and is indicated for patients with HBeAg+ or HBeAg– chronic hepatitis B (14-16). It is very well tolerated with few adverse effects. After one year of therapy, studies (17-19) have found a loss of serum HBV DNA and HBeAg in approximately 44% and 30% of HBeAg+ patients, respectively. In HBeAg– patients, loss of HBV DNA was seen in approximately 39% of patients (20). At the end of therapy, 53% of HBeAg+ patients and 60% of HBeAg– patients demonstrated histological improvement on liver biopsy. Among patients who experienced HBeAg seroconversion during treatment, the durability of response after cessation ranged from 38% to 77% (2). The major concern with lamivudine is the development of resistance, which increases with the duration of therapy. Long-term studies (2, 14) have now shown a resistance rate of 70% at four years and the benefit of the drug is lost. Lamivudine resistance can be associated with acute exacerbations of liver disease and, in rare cases, hepatic decompensation. For patients with confirmed resistance, the options include continuing the treatment as long as a benefit can be shown, discontinuing and monitoring for flares, or switching to other agents (2). A concern is that the development of resistance to lamivudine may compromise the response to other antiviral agents, particularly telbivudine and entecavir (2, 14). Although much less costly than other antivirals, lamivudine may no longer be the most effective first-line choice antiviral agent for chronic HBV infection, except in circumstances in which resistance development may be minimized (14).

Adefovir, a purine nucleotide analogue, is effective for treatment of HBeAg+, HBeAg– and lamivudine-refractory chronic HBV. It is less potent as an antiviral agent, but is associated with less resistance development. The optimal treatment duration for patients with chronic hepatitis B is unknown. In one study (21), loss of serum HBV DNA and HBeAg occurred in 21% and 24% of HBeAg+ patients, respectively, after 48 weeks. HBeAg seroconversion after one year is approximately 12%, but may increase to 40% over time (22). Loss of HBV DNA occurred in 52% of HBeAg– patients (23). Overall, 53% of HBeAg+ patients and 64% of HBeAg–
patients showed improvement on liver biopsy histology. Adefovir is generally well tolerated, but it has been associated with renal dysfunction and hypophosphatemia, although the latter does not appear to have clinical consequences. A potential advantage of adefovir for treatment of chronic hepatitis B is the absence of resistance after one year of therapy and only 18% resistance after four years of therapy (23). Adefovir may be a valuable choice for patients who are refractory to lamivudine due to resistance development (2,14).

Telbivudine is a pyrimidine L-nucleoside analogue with potent activity against HBV; in a large, multicentre trial, telbivudine was associated with similar rates of loss of HBeAg at one year (26%) compared with lamivudine (23%), but with higher rates of loss (60% versus 40%) of detectable HBV DNA (24). Although rates of loss of HBeAg were still similar after two years, resistance with breakthrough was found in 22% versus 35% of HBeAg+ patients and 9% versus 22% of HBeAg- patients receiving telbivudine versus lamivudine. Although telbivudine may suppress viral replication to a greater degree than lamivudine, and rates of resistance are lower with than lamivudine, resistance rates are higher than with other approved therapies. Telbivudine is generally well tolerated, but an increase in creatinine kinase has been reported in approximately 12% of patients; occasional cases of symptomatic myositis have also been reported (24).

Entecavir is an acyclic guanosine analogue with potent activity against HBV and has activity against both wild-type and lamivudine-resistant HBV. Three multinational, randomized, double-blind, pivotal, phase III entecavir safety and efficacy trials (25-27) have been conducted to date, and entecavir has demonstrated excellent potency, high rates of suppression of HBV DNA levels and improvements in liver histology. Rates of clearance of HBeAg (21% at one year and 39% at three years) were similar to those of other nucleoside agents, and resistance occurred in less than 1% of nucleosideneative patients after one- and two-year courses of therapy (28). Unfortunately, entecavir resistance has been found in over 30% of patients with pre-existing lamivudine resistance, and the latter appears to predispose to entecavir resistance (29).

Longer trial extensions involving lamivudine-refractory patients are needed to determine entecavir's optimal duration, long-term safety and durability of response, including rate of resistance. Entecavir is generally well tolerated. Recent guidelines (14) suggest that entecavir should be used in nucleosideneative patients and in lamivudine-refractory patients when no other alternative is available.

New Canadian consensus guidelines (14) will provide physicians who treat patients with chronic HBV infection with a management scheme that reflects new developments in the literature and are a welcome addition. However, there remain many unanswered questions, including the cost-effectiveness of these new agents, the impact of major shifts in initial therapy on provincial budgets, the optimal choices in the setting of multiple antiviral drug resistance, optimal timing of discontinuation of antivirals, the role of combination therapy and the role of additional nucleoside analogues such as clevudine, elvucitabine, pradefovir and valtorcitabine. Although great strides have been made in the management of chronic HBV infection, there is a continuing need for additional well-designed trials and long-term cohort studies to provide evidence to define optimal treatment for these patients.

REFERENCES

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