**Hepatitis B viral mutants and their relevance to the Canadian health care system**

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Over the past 10 years, an increasing number of mutations in the hepatitis B virus genome have been described. While the majority of these mutations appear to be 'silent' or not clinically relevant, some have been described in association with increased severity of disease (core and basic core promoter mutations), evasion of immunological surveillance (S escape mutants), hepatocellular carcinogenesis (X mutants) and resistance to antiviral agents (DNA polymerase mutations). The molecular events and the clinical consequences thereof are reviewed.

**Key Words:** Hepatitis B virus

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There are few areas of medicine that can generate as much confusion and despair in the minds of medical practitioners as hepatitis B viral (HBV) infections. A large part of that response stems from the array of serological tests associated with the virus and, to some extent, the varying interpretation of the results in the context of the clinical scenario and other findings. Thus, it was with much relief that physicians learned in the early 1980s of the development of a vaccine for HBV that held promise to eradicate the virus and thereby render the need to understand HBV only for historical interest. More recently, successes with antiviral therapies in the 1990s have reinforced that initial optimism. Did both advances provide the medical community with false hopes? Are the days for this virus truly numbered, or will physicians continue to manage HBV-infected individuals for generations to come? Unfortunately, the ever-increasing number of mutations described in response to immunoprophylaxis and antiviral therapies raises the possibility that HBV will not disappear but will continue to cause significant morbidity and mortality in a large percentage of the world’s population for at least the foreseeable future.

The principal purpose of this review is to provide medical practitioners with an account of the current situation in the battle against HBV and, specifically, to determine whether there is need for concern regarding the mutations
or ‘counterattack’ the virus has developed in its struggle for survival. However, any such account must begin with an attempt to put the clinical problem in its proper perspective, a working understanding of the replicative cycle of the wild-type (nonmutated) virus and an outline of our present interpretations of HBV serology.

CLINICAL PROBLEM
The prevalence of HBV varies between 0.1% and 15% in different parts of the world (1). Because prevalence rates are highest in areas where the world’s population is greatest (eg, Southeast Asia), the overall prevalence has been estimated to be approximately 5%, or 350 million individuals. Somewhat different concerns exist regarding the natural history of HBV infections in children and adults. Whereas their HBV infections tend to be mild and often subclinical, children’s propensity to develop chronic HBV carrier states (infections that persist beyond six months) is high (90% in newborns and 15% to 30% in adolescents). On the other hand, infected adults more often develop clinically apparent hepatitis and even fulminant hepatic failure, but much less often (5% to 10% of cases) develop chronic carrier states. Many but not all chronic HBV carriers (approximately 20% to 30%) eventually die of liver failure and/or hepatocellular carcinoma (HCC). Those at the greatest risk of such outcomes are males, those infected early in life and those residing in areas of the world where exposure to cocarcinogens such as aflatoxins is common; there is a 40% to 50% chance the deaths of people in these groups will result directly or indirectly from the consequences of HBV infection (2). While recent advances in immunoprophylaxis and antiviral therapies are expected to decrease this figure significantly, the introduction of such measures to impoverished regions where their need is greatest is a significant challenge to the health care delivery system of those nations.

HBV STRUCTURE
HBV is a small (42 nm), double-stranded DNA virus belonging to a family of hepatitis DNA viruses called Hepadnaviridae (3). Other well known members of the family include the woodchuck, ground squirrel and duck hepatitis viruses. Although genetically similar, these viruses cannot establish infections in humans. While attempts have been made to extrapolate the natural history of these viruses from infected animals to humans, the very different immune response of the human, combined with the fact that HBV is not inherently virulent but rather induces inflammation and injury through immune-mediated mechanisms, renders such extrapolations tenuous at best. The same argument can be made, although it rarely is, against their use as models for antiviral therapy.

Structurally, the HBV consists of an outer lipoprotein envelope and an inner protein/DNA nucleocapsid. The lipid component of the envelope is derived from the host cell, while the virus itself synthesizes the protein component (4). The protein consists of three related proteins referred to as large (L), middle (M) and small (S) hepatitis B surface antigens (HBsAg). Sequencing of the genes responsible for encoding these proteins has resulted in a relatively new classification system for HBV termed ‘HBV genotyping’. To date, six genotypes (A to F) with epidemiological but not clinical relevance have been identified. Genotype A is most common in North America.

Each HBsAg protein (L, M and S) possesses certain functions and properties that are important, if not essential, for viral survival. For example, LHBsAg is thought to contain the binding site required for viral entry into the host hepatocyte (5,6). It is also essential for viral assembly (7). When intracellular levels of LHBsAg are excessive, secretion of intact virus from cells is altered (4). Extremely high levels have been associated with the development of HCC in mice (8). LHBsAg also contains B and T cell epitopes that play a role in the humoral and cellular response to infection (9). Finally, a smaller or ‘truncated’ version of LHBsAg has the ability to stimulate mRNA synthesis and serve as a 'transcriptional activator', which might be relevant to hepatocarcinogenesis (8).

Fewer functional roles have been identified for MHBsAg. It is clearly not essential for viral assembly, but mutations at the site responsible for its own transcription, the initiation codon, have been described in Mediterranean patients with fulminant hepatic failure and HCC (10-12).

The smallest HBsAg, SHBsAg, is the predominant component of the viral envelope. It contains a hydrophilic region between amino acids 100 and 160 that results in a double-looped structure being displayed on the outer surface of the virus (13). The second loop between amino acids 124 and 147 is conserved in all HBV genotypes and serves as the target for neutralizing antibodies (antibody to HBsAg [anti-HBs]), be they naturally acquired, a result of passive prophylaxis with hepatitis B immunoglobulin (HBIG) or in response to active prophylaxis with HBV vaccine (natural and recombinant) (14).

Encompassed by the viral envelope is the nucleocapsid or core of the virus. The core consists of approximately 200 protein particles that together constitute the hepatitis B core antigen (HBeAg). Each core particle in turn consists of a large (185 amino acid) and a small (29 amino acid) segment (15).

In addition to capsid formation, core particles are responsible for packaging the mRNA required for viral replication, together with the enzyme involved in the replicative process (16). Other functions of core particles include trafficking of the capsid in the host cell, containment within the viral envelope, and provision of B and T cell epitopes that render the core immunogenic to the host (17).

A slightly larger version of core particles is the hepatitis B e antigen (HBeAg). Whereas the core protein is the translation product of the C gene, translation from an upstream initiation codon results in a precore protein, which is post-translationally modified at both the C and N terminal ends to form the soluble HBeAg. This 17 kD secretory protein is synthesized in concert with viral repli-
eral cellular genes that regulate cell growth (eg, c-variety of properties, including the ability to activate sev-
during the natural history of HBV infection, possesses a
protein. This protein, which has a low level of expression
the viral replicative cycle (20).
domain is the RNase H domain (residues 681 to 832),
domain with five conserved motifs (A to E). The fourth
reverse transcriptase domain in turn contains a catalytic
RNA- and DNA-dependent polymerase activity (20). This
scriptase (residues 337 to 630) domain that possesses both
separate the priming domain from the third, reverse tran-
consisting of 153 amino acids (residues 179 to 336) that
plate, and the second is a nonfunctional ‘spacer’ domain
The first domain (residues 1 to 173) primes the viral tem-
(32,33).
More recently, HBX has been reported to inhibit cellular
active in the nucleus, where it directly interacts with the
chemical carcinogens and accelerates c-
induces HCC in transgenic mice (27). It should be noted,
however, that in some mice exposed to HBX, no pathology
occurs, or HBX merely increases their susceptibility to
HBcAg in the liver toward an innocuous circulating pro-
tin and, therefore, serves as a surrogate marker of viral
replicative activity (18). It serves the virus by acting as an
immune tolerogen, thereby diverting host responses from
HBcAg in the liver toward an innocuous circulating pro-
tin. The only structural difference between core and e pro-
teins is that e proteins lack the arginine-rich, 35-amino acid
end of the core particle (17). Despite the rather subtle
structural difference, it is sufficient to result in two very dif-
ferent host responses; HBcAg induces a T helper 2 lympho-
cyte response, while HBcAg elicits a T helper 1 lymphocyte
response and cellular immunity (19).
Although not a structural component of the intact virus,
HBV replication is dependent on the synthesis of a unique
832-amino acid enzyme known as DNA polymerase
domain consists of four functional domains. The first domain (residues 1 to 173) primes the viral tem-
and the second is a nonfunctional ‘spacer’ domain
consisting of 153 amino acids (residues 179 to 336) that
separate the priming domain from the third, reverse tran-
scriptase (residues 337 to 630) domain that possesses both
RNA- and DNA-dependent polymerase activity (20). This
reverse transcriptase domain in turn contains a catalytic
domain with five conserved motifs (A to E). The fourth
domain is the RNase H domain (residues 681 to 832),
which serves to destroy residual viral mRNA resulting from the
viral replicative cycle (20).
The smallest HBV protein is the 154-amino acid HBX
protein. This protein, which has a low level of expression
during the natural history of HBV infection, possesses a
variety of properties, including the ability to activate sev-
eral cellular genes that regulate cell growth (eg, c-jun, c-fos,
c-myc, class II and III promoters), induce cell proliferation
in quiescent fibroblasts, reduce clonal outgrowth of cells,
increase apoptosis (via a p53-dependent pathway) and
increase a late G1 cell cycle block (21-26). Moreover, HBX
not only transforms certain cell types in vitro, but also
induces HCC in transgenic mice (27). It should be noted,
however, that in some mice exposed to HBX, no pathology
occurs, or HBX merely increases their susceptibility to
chemical carcinogens and accelerates c-myc-induced HCC
but does not act independently of these factors (28,29).
The majority of HBX is cytosolic, but a small portion is
active in the nucleus, where it directly interacts with the
P53 tumour suppressor and DNA repair factor (30,31).
More recently, HBX has been reported to inhibit cellular
protein degradation via a serine protease inhibitor domain
(32,33).

HBV REPLICATIVE CYCLE
Many questions remain unresolved with respect to the pre-
cise mechanism whereby HBV viral protein synthesis and
replication occur. For example, the host cell receptor
responsible for HBV entry has yet to be identified, as does
how the virus disassembles and translocates to the host
nucleus. Once within the nucleus, however, certain events
are known to occur (34). One of the earliest is the comple-
ton of the incomplete DNA strand that constitutes the
double-stranded DNA structure of the viral genome. This
event is mediated by a cellular DNA-dependent, RNA
polymerase enzyme. The end result is a closed, circular,
covalently coiled DNA that is now capable of mRNA syn-
thesis.
Four mRNAs are encoded by the virus. The largest is a
3.5 kb mRNA known as pregenomic HBV-RNA (pgRNA).
Synthesis of this mRNA is under the regulation of a region
of the viral genome called the basic core promoter (BCP).
pgRNA serves as the template for both HBcAg and DNA-P
synthesis. Another form of the 3.5 kb transcript – preco-
 mrNA – is also under the control of BCP and is responsi-
ble for HBeAg synthesis (35). LHBsAg is derived from a
2.4 kb pre-S1 mRNA, while two smaller 2.1 kb mRNAs are
responsible for MHBsAg and SHBsAg synthesis, respec-
tively. To date, the 0.7 kb HBX mRNA has only been
described in transfected cell lines (36,37).

When sufficient intracellular levels of all four mRNAs
are reached, viral assembly occurs. Within the developing
virion, DNA-P binds to pregenomic mRNA and is translo-
cated to the encapsidation signal epsilon, and reverse tran-
scription begins (38). A negative-strand DNA is produced,
while RNAse H of the polymerase protein destroys much of
the pgRNA (15 to 18 nucleotides remain to serve as a
primer for the subsequent synthesis of the positive DNA
strand) (39). The assembled nucleocapsid is then
enveloped by the lipoprotein membrane formed from the
cell lipid bilayer and viral surface proteins. All but a small
amount of the nucleocapsid material required to maintain
viral DNA within the host nucleus is exported through an
unidentified secretory pathway (40).

HBV SEROLOGY
The following are the six serological tests most commonly
used in clinical practice.

HBsAg
Historically, the first serological marker to be described in
association with HBV infections was HBsAg, or ‘Australia
antigen’. The presence of this marker in the blood generally
indicates the presence of an HBV infection. As discussed
above, HBsAg is synthesized and secreted by infected hepa-
tocytes in excess (1000- to 10,000-fold) of the numbers of
actual virions produced by the liver; therefore, while its
detection indicates the presence of an HBV infection, lev-
els of HBsAg do not necessarily correlate with the extent of
viral replication (18). False positive HBsAg results are rare
and are largely limited to a small subset of vaccine recipi-
ents (typically newborns and dialysis patients) who, for
short periods of time after vaccination, may be HBsAg pos-
itive but HBV-DNA negative (41). False negative results
most commonly occur during the so-called ‘window’ phase
of acute HBV infections, when HBsAg levels are below the
detection limits of most assays and antibody to HBsAg has
yet to appear. Because individuals in this phase of infection
may still be HBV-DNA positive, they should continue to be
considered infected and, therefore, potentially infectious to

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others. The window phase of HBV infections can last for weeks and, rarely, years (42). Throughout this period, antibody to HBeAg (anti-HBe) may serve as the sole indicator of recent HBV infection. Negative HBsAg results, despite the presence of infection, can also occur in HBV-vaccinated infants and in transplant recipients treated with high doses of hepatitis B immunoglobulins (see below).

Anti-HBs
Just as the presence of HBsAg in the blood generally indicates an HBV infection, anti-HBs indicates immunity to HBV. Whether that immunity was obtained as a result of recovery from a natural infection, or as a response to HBsAg vaccination or the recent (past three to six months) administration of hepatitis B immunoglobulin can usually be ascertained by questioning the patient and additional serological testing. At some time during chronic HBV infection, approximately 30% of HBsAg-positive carriers will have concurrent anti-HBs (43). The anti-HBs in this setting is often present in low titres and is heterotypic, ie, directed toward an HBV subtype other than that in the infected host (44).

Anti-HBc
Although the 200 core peptides that make up the nucleocapsid of the intact virion are not secreted by hepatocytes in any form other than that encompassed by HBsAg (ie, the intact virion), core proteins referred to as HBeAg are integrated into hepatocyte cell membranes and thereby induce an anti-HBc response. This response is prompt, and lasts throughout the course of infection and for decades beyond clearance of the virus (18). Thus, the presence of anti-HBc in the circulation indicates that an individual has been infected with HBV sometime in their life. Whether the infection is ongoing or has resolved is determined by the presence or absence of HBsAg and anti-HBs, respectively. The presence of immunoglobulin M anti-HBc generally indicates that infection occurred within the past three to six months; however, positive immunoglobulin M anti-HBc results can also occur during severe exacerbations of chronic HBV infections (45). False positive and negative anti-HBc results are uncommon (less than 5% of cases) with modern serological assays.

HBeAg
An additional antigen used in HBV serological testing is HBeAg. As discussed above, HBeAg consists of a large amino acid peptide that is synthesized and secreted by hepatocytes in proportion to the extent of viral replication. Thus, the presence and quantity of HBeAg in the circulation reflect the level of viral replication and, therefore, the degree of infectivity to others (46).

Antibody to HBeAg
Other than in the setting of precore mutant infections, the presence of antibody to HBeAg (anti-HBe) generally indicates low levels of HBV replication and low degrees of infectivity (47). Both HBeAg and anti-HBe may be absent during the transition period between active and inactive viral replicative states (47). Concurrent HBeAg and anti-HBe positivity is rare but can be seen during the same transition period.

HBV-DNA
HBV-DNA testing by relatively insensitive assays (non-polymerase chain reaction [PCR]-based techniques) is increasingly being used to confirm the presence of HBV infection, determine levels of viral replication/load and predict responses to antiviral agents (48). The clinical relevance of a PCR-positive HBV-DNA test in an HBsAg-negative, anti-HBs-positive individual has yet to be determined.

HBV MUTANTS
The half-life of HBV in the blood has been estimated to be one day (49). As HBV-DNA levels tend to be relatively constant, given an average viral load, approximately 10^{11} HBV particles must be released into the circulation daily (49). Thus, considerable viral replication is presumed to be underway within the liver. With each replicative cycle, the opportunity exists for mutations to occur in the viral genome. Why then, at such a high rate of replicative activity, have more HBV mutants not been recognized? The answer to this important question is almost certainly multifactorial. Two frequently cited explanations follow.

First, many additional mutants exist but have yet to be identified. Indeed, rapid advances in molecular biological techniques have confirmed that a large number of HBV mutants do exist. Moreover, many of these mutants are capable of inducing acute and chronic liver diseases that were previously ascribed to other known or suspected hepatotropic viruses. It should be noted that there are two commonly employed strategies for detecting viral mutants:

1. PCR amplification of the viral genome followed by cloning of the PCR product, selection and sequencing of individual clones, and

2. PCR amplification and direct sequencing of the PCR product.

The former cloning strategy provides sequences of only one strain (dominant or not), whereas the latter direct strategy provides a more complete analysis of the overall viral population, although only of the predominant strains (34). Neither strategy provides extensive information about HBV quasispecies, with the clonal assay being limited by the number of clones selected for evaluation. Also relevant to mutation detection is the cross-sectional nature of many studies, wherein the emergence of HBV mutants would not be detected unless a longitudinal study format were in place. Indeed, the distinction between ‘mutations’ and ‘variations’ is based on the chronology of the viral form, with mutations being considered nucleotide changes arising in an evolutionary fashion over time and within the same source, whereas variations refer to nucleotide differences.
from the 23 fully sequenced HBV genomes registered in GenBank that are stable over time and within sources (34).

The second, perhaps more relevant, explanation for the perceived paucity of HBV mutants relates to the unique structure of the HBV genome. The compact nature of the genome (Figure 1) defines four overlapping open reading frames (ORF). These ORFs encode the envelope (pre-s/s), core (pre-c/c), polymerase and X proteins. The extent of the overlap is such that more than 50% of the genome is translated by more than one ORF, and every nucleotide is located within at least one coding region (15). Thus, a single mutation in one ORF (eg, pre-s/s) will affect the structure of the overlapping gene product (polymerase enzyme). In many instances, this domino effect renders the virus either replicatively incompetent or incapable of infecting additional hepatocytes (15). In many cases, mutant viruses may also have a reduced fitness, making them inefficient competitors for the wild-type virus. Additional changes may alter immune recognition and clearance. There are numerous possible additional effects. Thus, the dynamics of both replication and clearance are key in evaluating mutant emergence in patients.

On the other hand, there are features of the virus and its replicative cycle that encourage mutation development. One such feature is the lack of proofreading capabilities of the HBV-DNA-P enzyme (50). Proofreading is a mechanism whereby mutations – whether they be point (nucleotide substitutions), insertional or deletional (which cause fatal frame shifts) – are identified by replicative enzymes and either replication ceases or a dysfunctional virus is produced. Unlike RNA viruses, most DNA viruses incorporate cell enzymes that have proofreading capacity; therefore, in general, fewer mutations are tolerated with DNA viral infections. HBV, which uses its own reverse transcriptase/DNA polymerase enzyme, is an exception. The end result is that approximately one in $10^4$ nucleotide base misincorporations occur per year (51). If these mutations offered survival advantages over the wild-type virus, such as greater replicative activity, the ability to evade the host’s immune response or resistance to antiviral agents, the mutants would persist and might become the predominant viral pathogen of the infected liver.

**S mutants**

The target of the host’s humoral response to HBV is the hydrophilic region of HBsAg between amino acid residues 100 and 160, the so-called ‘a’ epitope. Thus, a distinct survival advantage would be afforded to HBV variants with mutation(s) in this (more precisely between amino acid residues 124 and 147) or adjacent regions that interfere with the binding of neutralizing anti-HBs. Because the ‘a’ epitope domain is within the RT domain of the DNA-P (whereas pre-S1/S2 domains correspond with the spacer domain of DNA-P), advantageous changes in viral replicative activity might also occur with mutations in this region. Such a mutation was first described in an Italian child who developed an HBV infection, despite having received complete vaccination and HBIG at birth (52). The mutation resulted in a glycine to arginine switch at amino acid 145. Other common mutations reported in this region include aspartate-144-alanine, methionine-133-leucine, glutamine-129-histidine and isoleucine/threonine-126-alanine (53). These mutations have been described in many areas of the world but appear to be most common in Asian infants; in this group, 2% to 3% of vaccine recipients at birth were found to have glycine to arginine mutations at this site (53). An increased risk of developing S mutations has been described in infants of mothers with high viral loads and mutations elsewhere in the mother’s HBV S gene (54). Glycine to arginine mutations are also common in liver transplant recipients receiving HBIG (53). Less frequently, they develop spontaneously during the course of a chronic HBV infection (55,56).

Some of these mutations, which are often referred to as ‘S escape mutants’, have been described in association with important clinical events. For example, the threonine-126-alanine mutation has been identified in vaccinated infants who subsequently developed fulminant hepatic failure (57). Of more diagnostic concern are nucleotide insertions in the area of amino acids 121 to 124, which can result in false negative HBsAg test results and thereby represent a risk to the health of an undiagnosed patient and the safety of the blood transfusion system (55-58).

The study that has most extensively documented the prevalence of S escape mutants is that reported by Hsu et al (59). Investigators studied approximately 1500 stored sera sampled from children just before universal HBV vaccination in their country (1984), and from similar sized groups of children belonging to the same age groups five and 10 years after the introduction of universal vaccination. The results of the study revealed that the contribution of S escape mutants to the overall prevalence of vaccine failures (HBV-
DNA-positive children) increased from 7.8% in 1984 to 19.6% in 1989 to 23.1% in 1994. That vaccination was responsible for the increases was supported by the finding that the mutation rate was 36.4% in vaccinated children but only 9.9% in unvaccinated children. Moreover, those with S escape mutants were younger than those without, suggesting that vaccine-induced pressure resulted in a more rapid emergence of the mutant genotype. Finally, mutations in the neutralizing region were more common in the 10-year postuniversal vaccination cohort. Of interest, the absolute numbers of carriers with mutants did not increase over the 10 years of the study (eight of 1200, nine of 1134 and five of 1515 in the years 1984, 1989 and 1994, respectively), suggesting that, although these mutants constitute an increasing percentage of vaccine failures, they are not becoming more widespread.

The clinical concerns associated with S escape mutants lie in the following four areas.

- Although uncommon, their failure to be detected by commercially available tests for HBsAg raises the possibility that carriers of these mutants could enter the blood donor population (60). In this regard, it is important to note that, in more than 95% of cases, anti-HBc is and remains strongly positive (59). Other diagnostic tests that can be positive in this setting are HBV-DNA and HBeAg (59).

- Despite encouraging results in vaccinated chimpanzees, HBIG and HBV vaccination does not provide universal or complete protection from infection with these mutants in humans (61). Fortunately, intrafamilial spread of the mutants has yet to be described.

- Like other HBV forms, these mutants have the capacity to induce both acute and chronic liver disease, as well as HCC (62,63).

- Because mothers of children with S escape mutants need not be HBsAg and/or and anti-HBc positive, and need not have mutants themselves (indeed, more often do not), horizontal as well as vertical transmission appears likely (60).

**Precore and core promoter mutants**

More mutations of the viral genome may be tolerated in the large region of the precore/core ORF, which is not overlapped by another ORF. The most common and extensively studied of these is the glycine to arginine point mutation at nucleotide 1762 and a glycine nucleotide substitution at nucleotide 1764. The status of the 1858 nucleotide site may be clinically important in its own right because the predominant genotype in North America and Europe is A, which has a cytosine at position 1858 rather than the thymidine nucleotide required to stabilize the arginine base at the corresponding 1896 site (66). The same low prevalence would also be predicted in regions where the F genotype is common, where a thymidine is also lacking at nucleotide 1858. The status of the 1858 nucleotide site may be clinically important in its own right because studies have demonstrated that patients without mutations at position 1858 had more benign histological findings (67). It should be noted that not all HBeAg-negative, HBV-DNA-positive patients have the precore mutation at position 1858. A small percentage, estimated at approximately 10% to 15%, have other mutations, insertions and/or deletions, resulting in in-frame shifts that explain their HBeAg negativity (69). Glycine to arginine mutations at nucleotides 1898 and 1899 have also been reported to be associated with the severity of liver disease (70,71).

Although initially described in association with histologically active liver disease, including fulminant hepatic failure, the precore mutant has more recently been identified in chronic HBV carriers without biochemical or histological evidence of liver disease (67). Indeed, in a recent population-based study of Canadian Inuit, none of the approximately 35 precore mutant carriers identified had clinical or biochemical evidence of liver disease (72). Moreover, there are data indicating that reactivation of disease is not associated with the emergence of precore mutations and that wild-type virus is associated with more inflammation and fibrosis than precore mutants (73). Finally, HBV-DNA levels are not increased in patients with precore mutant infections compared with those with wild-type infection (74,75). Thus, the mutation itself is not uniformly pathogenic. These findings have led to searches for comutations that might explain the more virulent forms of precore infections originally described. Strong candidates are point mutations and short deletions or insertions in the core promoter/enhancer II (nucleotide 1634 to 1782) region, which not only limit the transcription of precore mRNA and thereby HBeAg synthesis, but also enhances transcription of core mRNA and with it, HBeAg and polymerase enzyme synthesis (76). The most frequently described mutations in this region are an arginine to tyrosine nucleotide substitution at nucleotide 1762 and a glycine to arginine substitution at nucleotide 1764. The existence of mutations at these sites correlates with disease...
progression and has been implicated in hepatocarcinogene-
sis (77-79). Clinical studies have shown that 30% of pre-
sumably healthy HBV carriers attempting to donate blood
had core promoter mutations compared with 90% to 100% of
patients with active disease, cirrhosis or HCC (80). This
increased pathogenicity presumably relates to increased
core protein synthesis and viral replication, and decreases
the synthesis of the immunotolerogen HBeAg. However,
not all studies have identified an association between
1762/1764 mutations and the severity of liver disease (81).
Whether precore and/or core promoter mutations predis-
pose the patient to a rapidly progressive fibrosing, cholesta-
tic form of liver disease after liver transplantation is unclear
(82-84).

In terms of therapeutic implications, initial reports sug-
gested that, if more than 20% of the viral population within
a patient consists of precore mutants, then a poor response
to interferon therapy can be expected (85). On the other
hand, more recent data suggest that a high prevalence of
precore mutants results in earlier responses to interferon
therapy (86). What is clear is that relapses are more com-
mon when interferon therapy is withdrawn; however, this
may be also be related to associated mutations in the core
promoter or core region rather than in the precore region
(87). Indeed, the importance of the core promoter region in
terms of predicting the response to interferon therapy has
recently been emphasized in a report describing that
responses can be predicted in approximately 80% of HBeAg-
positive and -negative cases by documenting the presence
or absence, respectively, of the core promoter mutation at
position 1764 (88). Similar findings (rapid initial responses
but high relapse rates in patients with precore mutants) have
also been described with nucleoside analogue therapy (89).

HBCAg mutants
The most immunodominant epitopes of HBCAg are
between amino acids 50 and 69, and 61 and 85 (90).
Mutations in this region have been described in HBV
patients with active hepatitis but not during the immune-
tolerant phase of the infection (91,92). If the mutations,
which likely occur as a result of B and T cell pressure on the
core antigen, are present before interferon therapy, response
to treatment is less likely (88,93). Deletions in the core
region have also been described and have been shown to
result in decreased cytotoxic T cell responses and viral repli-
cation (94). Similar deletions may play a role in converting
an immune-tolerant to an immune-intolerant state and in
the progression of acute to chronic HBV infection (95).

DNA-P mutants
Mutations in the catalytic domain of DNA-P are not
detected normally but have been described in association
with nucleoside analogue therapy (96). In the case of
lamivudine, the first nucleoside analogue licensed for the
treatment of HBV, a methionine-552-isoleucine or methio-
nine-552-valine mutation in the conserved tyrosine-methio-
nine-aspartate-aspartate' (YMDD) motif that comprises part
of the active site (domain C) of the reverse transcriptase
has been described, often in association with an upstream
leucine-528-methionine substitution (97). Famciclovir,
another nucleoside analogue with anti-HBV properties, can
induce valine-521-leucine and leucine-528-methionine
mutations within the B domain of DNA-P (98). These
mutations decrease but do not eliminate the affinity of
nucleoside analogues for the DNA-P enzyme. As a result,
with the development of these mutations, viral replication
increases, but not to levels documented before initiation of
treatment (99-101). Because the mutant virus is replica-
tively compromised in the vital region of DNA-P activity,
wild-type virus returns on cessation of drug therapy (102,103).

Of interest, the ‘a’ determinant is located in the variable
linker region between the A (410 to 426) and B (498 to
528) domains of DNA-P. Thus, while YMDD mutations
could result in HBsAg changes, they are unlikely to do so in
the critical antigenic site of HBsAg. Nonetheless, patients
with both YMDD and S escape mutations have been
described (104).

HBX mutations
Data are only just emerging on mutations in the X ORF,
which overlap with precore stop codon mutants and BCP
mutants. Preliminary findings describe a novel class of HBX
mutants in Asian patients with HCC (105). These muta-
tions were found to negate X-induced inhibition of clonal
outgrowth and increase apoptosis. Thus, X mutants
increased clonal outgrowth and decreased apoptosis, sug-
gesting a possible carcinogenic role. While this is an
intriguing hypothesis, it should be noted the HCC tissue
sources for the HBX mutations detected also contained
HBsAg mutations, which may have been relevant to HCC
development (106).

CONCLUSIONS
Available data allow the following conclusions to be made
regarding the S escape mutant.
• The mutants constitute an increasing percentage of
  ‘vaccine failures’, but their prevalence in the general
  population does not appear to be increasing.
• The overall prevalence in HBV-vaccinated Southeast
  Asian children born of HBV carrier mothers appears
  to be low (1% to 2%).
• Most often, these mutations are associated with the
  following serological profile: HBsAg, anti-HBs
  positive and anti-HBc positive. However, they can
  also exist as HBsAg negative, anti-HBs positive and
  anti-HBc positive or anti-HBc positive alone.
• S escape mutants can be associated with active viral
  replication (approximately 50% are HBeAg positive).
• Although precise figures are not yet available, these
  mutants can be associated with acute and chronic
  liver disease.
• Passive and active HBV immunity is unlikely to be
  protective against these mutants.
Regarding the precore/core promoter mutants:
- It is possible that the majority of precore mutants represent the natural evolution of chronic HBV infections.
- Worsening liver disease is more likely to be associated with core promoter rather than precore mutations.
- The natural history of acute precore/core promoter mutations is unclear.
- HBV-DNA levels tend to increase when core promoter mutations exist.

Regarding core mutants:
- Mutations to the core gene are less common than mutations in other areas of the viral genome.
- Core mutants appear to be associated with viral escape from immune surveillance and are relevant to the response (or lack thereof) to interferon therapy.

Regarding DNA-P mutations:
- DNA-P mutations are uncommon in the absence of nucleoside or nucleotide analogue therapy.
- The detection of these mutants in patients increases to approximately 50% after three years of continuous antiviral therapy.
- Although DNA-P mutations decrease the affinity of DNA-P for nucleoside/nucleotide analogue binding, a complete loss of antiviral effect is not observed.
- Finally, DNA-P mutations in the B and C catalytic domains are unlikely to significantly alter the dominant S epitope.

Regarding HBX mutants:
- Mutations in HBX may be associated with an increased risk of HCC.
- The same mutations hold promise as a marker of those HBV carriers who might be at risk of developing HCC in the future.

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