Two Core Promotor Mutations Identified in a Hepatitis B Virus Strain Associated with Fulminant Hepatitis Result in Enhanced Viral Replication

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Abstract

Viral mutations have been implicated in alteration of the biological phenotype of hepatitis B virus (HBV). We recently cloned and sequenced the viral genome of an HBV strain associated with an outbreak of fulminant hepatitis (FH strain). The FH strain contained numerous mutations in all genomic regions and was functionally characterized by a more efficient encapsidation of pregenomic RNA leading to highly enhanced replication. To define the responsible mutation(s) for the enhanced replication, we introduced individual mutations of the FH strain into a wild-type construct by oligonucleotide-directed mutagenesis. Analysis of viral replication showed that two adjacent mutations in the HBV core promoter (C to T at nucleotide 1768 and T to A at nucleotide 1770) led to high level replication. Similar to the FH strain, this mutant displayed the phenotype of enhanced encapsidation of pregenomic RNA. Functional studies in an encapsidation assay demonstrated that the identified mutations resulted in a minor increase of pregenomic RNA transcription (two- to threefold) and a major transcription-independent enhancement (> 10-fold) of viral encapsidation. Our results demonstrate that the two adjacent mutations in the HBV core promoter region are responsible for the enhanced replication of the FH strain. These two mutations, outside the previously described encapsidation signal, core, and polymerase polypeptides, appeared to affect a novel genetic element involved in viral encapsidation. (J. Clin. Invest. 1996, 98:2268–2276.) Key words: core promoter • encapsidation • HBX gene

Introduction

Hepatitis B virus (HBV)† is a partially double-stranded DNA virus that replicates through an RNA intermediate (for review, see references 1 and 2). The virally encoded reverse transcriptase-polymerase (RT-Pol) is essential for this unique form of genome replication. Viral replication occurs exclusively in the core particle which is assembled through complex interactions among pregenomic RNA, core protein, and RT-Pol in a process called encapsidation. The capsid containing the replicative intermediate is then enveloped by three forms of HBV surface antigens (HBsAg) in a lipid bilayer. Although the molecular mechanism of HBV replication has been largely elucidated with the identification of various essential elements (3–8), it is not clear whether any other sequences outside these elements play an important role in the replicative process.

We have previously described the association of an HBV variant with an outbreak of fulminant hepatitis (FH) B (9). This nosocomial outbreak of five cases was remarkable for an unusually severe course of acute HBV infection, leading to 100% mortality within a few days from hepatic and renal failure. The HBV strain (FH strain) responsible for this outbreak contained two G to A mutations in the precore open reading frame, one of them resulting in a stop codon (9). Cloning and sequencing of the entire genome of this viral strain revealed numerous mutations affecting conserved and nonconserved sequences in various regions of the HBV genome (10). Functional characterization of the FH strain in transfected hepatoma cells showed a highly increased viral replication (10). Enhanced replication of the FH strain was caused by a more efficient encapsidation of pregenomic RNA into the viral core particles. Our previous study demonstrated that the precore mutations were not responsible for enhanced replication (10). In the woodchuck model, the precore stop codon mutation had no effect on viral replication and the initial course of infection, but appeared to result in less chronic infection (11). These observations indicated that other mutation(s) in the HBV genome may be important for the altered biological behavior of the FH strain. In this study, we identified and characterized functionally the mutations responsible for the enhanced replication of this HBV strain.

Methods

Constructs. Replication-competent constructs of wild-type (adwR9, aywR9) and fulminant HBV (FHR9) strains were described previously (10). These constructs contained 1.2 times the genomic length of HBV. The HBX expression constructs HBXadw and HBXFH were obtained by subcloning the Hpal–BglII fragment of adwR9 and FHR9, respectively, into Smal and BamHI sites of pGEM7Z(+) (12). HBXMT5/6 was generated by ligating the XhoI–SacII fragment of FHR9, respectively, into SmaI and BamHI sites of pGEM7Z(+) (13). FHR9 was previously described (9). HBXMT5/6 was generated by ligating the XhoI–SacII fragment of FHR9, respectively, into SmaI and BamHI sites of pGEM7Z(+) (13). FHR9 was previously described (9).

1. Abbreviations used in this paper: ccc, covalently closed circular; FH, fulminant hepatitis; HBeAg, hepatitis B virus e antigen; HBsAg, hepatitis B virus surface antigen; HBV, hepatitis B virus; HBXKO, HBX knockout; hGH, human growth hormone; MT, mutant; nt, nucleotide; ORF, open reading frame; RT-Pol, reverse transcriptase-polymerase; WT, wild-type.
HBXadw and the SacI–BglII fragment of pSelectHBV mutant (MT)5/6 (described below) into the XhoI and BamHI sites of pGEM7Z (+). The core promoter constructs pXp2CPadw and pXp2CPMT5/6 were generated by digesting pHBXadw and HBXMT5/6 with AllIII (HBV nucleotide [nt] 1852), filling in the ends with Klenow, and then digesting with XhoI present in the 5' end of the multiple cloning site of pGEM7Z (+). The fragment was then cloned into the SalI and SmaI sites of pXp2 which contains a promoterless luciferase gene (12, 13). These constructs contained the luciferase gene driven by wild-type and mutant HBV core promoters including HBV sequences up to enhancer I (nt 965–1852). pCPadw lacZ was constructed by ligating the HincII (nt 1684–AllIII (nt 1852) fragment of pHBXadw, the AllIII-KpnI fragment of pE-lacZ (14) and the SmaI-KpnI fragment of pE-lacZ. pCPMT5/6lacZ was similarly constructed using pHBXMT5/6. In these two constructs, the CMV promoter of pE-lacZ was replaced with the wild-type or mutant HBV core promoter. All mutant constructs were confirmed by DNA sequencing. Covalently closed circular DNA was generated from wild-type (WT) (adwR9) or mutant (MT5/6R9) HBV-DNA as described previously (15).

**Oligonucleotide mutagenesis.** The Altered Sites In Vitro System (Promega Corp., Madison, WI) was used to introduce HBV core promoter, precore, core and HBX knockout (HBXKO) mutations: the first mutation with T to G at nucleotide 1848 (MT2), the second with a G to T and T to A at nucleotides 1754 and 1770 (MT5), the fourth with C to T at nucleotides 1768 and 1770 (MT5/6), and the fifth with T to A at nucleotide 1770 (MT6). The five mutant oligonucleotides were: 5'CTAATCATCTCGTGACATGTC3' (sense, nt 1837–1859) for MT2, 5'GGGGGAGGAGTATTAGGGTTAAGGTC3' (sense, nt 1744–1768) for MT3/4, 5'AGGTTAAGGTTAAGGAGG3' (sense, nt 1757–1781) for MT5, 5'AGGTTAAGGTTAAGGAGG3' (sense, nt 1757–1781) for MT5, and 5'AGGTTAAGGTTAAGGAGG3' (sense, nt 1757–1781) for MT6. To generate HBXKO mutations the following primers were used: 5'TGAAGCCGCAATAGATCTGGCCAA3' (to generate a stop codon for the HBX.2 protein p8) and 5'ACTCCCAGCAAGGTCAACGACCG3' (to generate a mutated start codon for the HBX.2 protein p8) and 5'AGGTTAAGGTTAAGGAGG3' (to generate a mutated start codon for the HBX.2 protein p8) and 5'ACTCCCAGCAAGGTCAACGACCG3' (to generate a mutated start codon for the HBX.2 protein p8) and 5'ACTCCCAGCAAGGTCAACGACCG3' (to generate a mutated start codon for the HBX.2 protein p8).

The plasmids pCPadw lacZ and pCPMT5/6 lacZ were generated by in vitro transcription of pLacZ (a generous gift from GE Healthcare Corp.). pHBVadw BH was constructed by subcloning an AatII (nt 1417)–BglII (nt 1986) fragment of HBVadw into pGEM7 (AatII and BamHI sites). Primer extension products and protected RNA fragments were analyzed on an 8% polyacrylamide-urea gel.

**Analysis of transfection efficiency.** The pTKGH plasmid containing the human growth hormone (hGH gene driven by the thymidine kinase enhancer and promoter) was cotransfected with various HBV constructs to control for transfection efficiency. Typically, 15 μg of HBV construct was cotransfected with 1 μg of pTKGH plasmid into HuH-7 cells grown in 10-cm dishes. From each transfection experiment, medium was harvested and hGH was measured with a radioimmunoassay from Nichols Institute Diagnostics (San Juan Capistrano, CA). A typical result for hGH is shown in Table I.

**Viral protein, RNA, and DNA analyses.** HBsAg and HBeAg were determined by commercially available radioimmunoassays (Abbott Laboratories, North Chicago, IL). 3 or 4 d after transfection, HuH-7 cells were harvested for viral RNA and DNA analysis. RNA was prepared by the guanidium isothiocyanate-acid-phenol method (16) analyzed by formaldehyde agarose gel electrophoresis (10 μg RNA) and hybridized with an HBV-specific probe. Viral replicative intermediates associated with intracellular core particles were isolated by ultracentrifugation of cell lysate through a 30% sucrose cushion and then analyzed by Southern blot hybridization (17).

**Analysis of core-associated RNA.** 3 d after transfection of HuH-7 cells with various HBV constructs, core particles were purified by ultracentrifugation as described above. The pellet containing the core particles was treated with Sphingolococcus aureus nuclease (Boehringer Mannheim, Indianapolis, IN) at a concentration of 8 μg/ml for 60 min at 37°C to eliminate any contaminating nucleic acids. Core-associated RNA was then isolated by the guanidium isothiocyanate-acid-phenol method twice (16). For primer extension analysis, an HBV primer (5'TCTAAGGCTCTCGATCACGACCTG3') spanning nt 2006–2030 in the antisense orientation was end labeled with γ-[32P]ATP and then reacted with RNA by a standard protocol (18). Core-associated RNA was compared with total RNA prepared as above. In addition, core-associated RNA was analyzed by RNAase-protection analysis (RPA II kit; Ambion Inc., Austin, TX). An α–[32P]PITC-labeled HBV-RNA probe (Aval–BglII fragment of HBVadw) was generated by in vitro transcription of the linearized (digested with Aval) plasmid pBVadvBH (In Vitro Transcription kit; Promega Corp.). pHBVadw BH was constructed by subcloning an AatII (nt 1417)–BglII (nt 1986) fragment of HBVadw into pGEM7 (AatII and BamHI sites). Primer extension products and protected RNA fragments were analyzed on an 8% polyacrylamide-urea gel.

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**Encapsidation assay.** *cis* and *trans* effects of MT5/6 on encapsidation of pregenomic RNA were studied using an encapsidation assay described previously (14, 19). Plasmid pE-LacZ (a generous gift from Dom Ganem) contains a fusion of the HBV encapsidation signal sequence (nt 1815–1986) and the lacZ gene driven by the CMV immediate-early promoter. Transcription from the CMV immediate-early promoter initiates at nt 1815 of the HBV sequence and leads to a chimeric HBVlacZ RNA. In plasmids pCPadw lacZ and pCPMT5/6 lacZ, transcription of the chimeric RNA is driven by the HBV core promoter (wild-type and mutant). By providing core and polymerase proteins in trans, the chimeric HBVlacZ RNA has been shown to be encapsidated into core particles in tissue culture (14). HBVlacZ RNA was analyzed by ribonuclease protection assay as described (5). Probe lacZ, which was generated by in vitro transcription of pLaZ (a generous gift from Dom Ganem, University of California, San Francisco, CA) is a 475-nt riboprobe with 50 polylinker nucleotides and 425 nt complementary to the lacZ sequence of the HBVlacZ RNA (5).

**Analysis of core promoter activity and transcription function of HBX.** Core promoter activities were studied by analyzing luciferase gene expression of the plasmids pXp2 CPadw and pXp2CPMT5/6 in transiently transfected HepG2 cells (0.5 μg DNA per six-well plate). The plasmids pXp2 (promoterless) and RSVLuc (containing the luciferase gene driven by the RSV promoter (12, 13) served as negative and positive controls. Luciferase activities were determined using the Luciferase reagent (Promega Corp.) with a luminometer. For analysis of HBX transactivation, 0.1 μg of HBXadw or HBXFH was cotransfected with 0.5 μg DNA of RSVLuc into triplicate of HepG2 cells maintained in six-well plates.
Results

Identification of mutation(s) responsible for enhanced replication. To further define the HBV sequences responsible for enhanced replication of FH strain, various regions of wild-type strain were replaced with equivalent regions of the FH strain in the replication competent construct adwR9. The chimeric constructs were analyzed for viral replication after transfection into HuH-7 cells (Fig. 1). The responsible region of the fulminant strain was mapped to a fragment spanning from nucleotide 1574 to 1986 (RsrII–BglII). This region contains nine mutations affecting conserved nucleotide sequences (10). We selected six of these mutations and introduced them individually or in combination into the wild-type adwR9 construct (MT2-7, Fig. 2). These mutations of the FH strain were located in the HBX gene, the core promoter, the encapsidation signal, and the core protein. These constructs were transfected into HuH-7 cells and viral replication was analyzed (Fig. 3). Only the construct MT5/6 demonstrated the phenotype of enhanced replication similar to the FH strain, while the other mutations had no effect. Densitometric quantification revealed an ~17-fold (17.5 ± 7.7, from four experiments) difference between wild-type and mutant (MT5/6) signal intensities for the replicative intermediates. The actual difference may be much greater because densitometry of autoradiography is only linear over one log. To examine whether both mutations 5 and 6 are required for the enhanced replication, constructs containing the individual mutations 5 and 6 (MT5 and MT6) were generated and analyzed for replication (Fig. 3). Mutants 5 or 6 by itself appeared to have only a minor nonsignificant effect on replication. Densitometric analysis from four experiments showed a 1.47 ± 0.63- and 1.49 ± 0.87-fold increase of replicative intermediates over the wild type for MT5 and MT6, respectively. To confirm the effect of MT5/6 on HBV replication in a more physiological setting, we generated covalently closed circular (ccc) DNA from wild-type and MT5/6 genome to better corre-

![Figure 1](image1.png)

**Figure 1.** Mapping of the HBV sequences responsible for enhanced replication. Two regions of adw strain, RsRII to XhoI (RX), RsRII to BglII (RBg), were replaced with equivalent regions of the FH strain in a replication-competent construct (adwR9). HuH-7 cells were transfected with the chimeric constructs and core particle–associated HBV DNAs were isolated and subjected to Southern blot analysis. Lane contents are as indicated.

![Figure 2](image2.png)

**Figure 2.** Construction of terminally redundant constructs of HBV mutants. (Top): The relative positions of RNA transcripts, ORFs, and regulatory elements (Enh 1 and 2) are displayed in reference to the genomic map of HBV. The wild-type and mutant constructs with terminal redundancy are shown below. The approximate position of the introduced mutations of the FH strain is shown by the respective numbers. (Bottom) Regional and numerical positions as well as nucleotide changes of mutations introduced by oligonucleotide-directed mutagenesis are shown.

![Figure 3](image3.png)

**Figure 3.** HBV replication in cells transfected with HBV constructs and ccc HBV-DNA. 4 d after transfection, HuH-7 cells were harvested and core particle–associated HBV DNAs were isolated and subjected to Southern analysis. Lane contents are as indicated. SS, single-stranded HBV DNA. Results are representative of four independent experiments.

<table>
<thead>
<tr>
<th>Mutant</th>
<th>HBV region</th>
<th>nt position</th>
<th>nt change</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>precore</td>
<td>1848</td>
<td>T to G</td>
</tr>
<tr>
<td>3</td>
<td>core promoter</td>
<td>1754</td>
<td>C to T</td>
</tr>
<tr>
<td>4</td>
<td>core promoter</td>
<td>1756</td>
<td>T to A</td>
</tr>
<tr>
<td>5</td>
<td>core promoter</td>
<td>1768</td>
<td>C to T</td>
</tr>
<tr>
<td>6</td>
<td>core promoter</td>
<td>1770</td>
<td>T to A</td>
</tr>
<tr>
<td>7</td>
<td>core</td>
<td>1982</td>
<td>T to C</td>
</tr>
</tbody>
</table>
spond to the natural template for the transcription of viral mRNAs. Analysis of viral replication of ccc HBV-DNA demonstrated a similar effect of MT5/6 on HBV replication as shown with the terminal-redundant R9 construct (Fig. 3). The two identified mutations (nt 1768 and 1770) are within the overlapping region of the HBV core promoter and the HBX gene.

Enhanced replication results from enhanced encapsidation of pregenomic RNA. To define the molecular mechanism of enhanced viral replication, we studied the functional effect of MT5/6 on viral encapsidation, transcription, protein expression, and function. Since the fulminant strain was characterized by a more efficient encapsidation of pregenomic RNA, we first analyzed the level of viral encapsidation of the mutant constructs in comparison to the wild-type strains (Fig. 4). Only construct MT5/6 showed a highly increased encapsidation of pregenomic RNA (Fig. 4). Densitometric analysis revealed that MT5/6 showed an ∼12-fold (12.1±4.4, from four experiments) increase in the signal intensity of encapsidated pregenomic RNA compared to the wild type. In contrast, the constructs containing the individual mutations MT5 or MT6 displayed only a minimal increase of encapsidation (MT5: 2.06±0.86-fold; MT6: 1.94±0.72-fold; from four experiments) compared with the wild-type constructs (Fig. 4). This finding was confirmed independently by ribonuclease protection analysis of encapsidated RNA (not shown).

The identified mutations affect core promoter activity leading to increased pregenomic RNA transcription. Since these two mutations were located within the core promoter, they might enhance transcription of pregenomic RNA resulting in the apparent increased encapsidation. Total viral RNAs from cells transfected with these constructs were analyzed by Northern blot and primer extension analysis. Terminal redundant R9 constructs containing the sequences of adw, ayw, MT5/6, MT5, MT6, and FH were transfected into HuH-7 cells. RNAs were purified and analyzed by Northern blot hybridization. Two species of HBV RNA, 3.5 and 2.4/2.1 kb, were present in all transfections (Fig. 5). In Fig. 5, densitometric quantitation showed a 2.4-fold increase of the MT5/6 3.5-kb RNA compared to the adw wild type when corrected for GAPDH analysis. The densitometric reading of 3.5-kb RNA for adw was 510 (arbitrary units) and for MT5/6 was 978; the densitometric reading of GAPDH for adw was 2335 and for MT5/6 was 1842. There was no significant difference in the 2.1- and 2.4-kb transcripts (Fig. 5). Although this twofold difference in 3.5-kb RNA level between WT adw and MT5/6 appeared small, it was confirmed by four other independent experiments. To study the effect of MT5/6 on pregenomic RNA transcription directly, primer extension analysis was performed. Primer extension of total viral RNA revealed three extension products: the shortest, representing the pregenomic RNA, and the two slightly longer species, representing the precore RNAs. The

Figure 4. Primer extension analysis of encapsidated RNA. 3 d after transfection, core-associated RNA was purified and subjected to primer extension analysis. Primer extension products were electrophoresed on an 8% polyacrylamide-urea gel and autoradiography using an intensifying screen with exposure time of 24 h at −80°C was performed. A sequencing reaction using the same primer on the adw strain is shown on the left (lanes marked A,G,C,T). Lanes are as indicated. Results are representative of four independent experiments.

Figure 5. HBV transcription in cells transfected with HBV constructs. 3 d after transfection, HuH-7 cells were harvested and their RNAs were isolated and subjected to Northern blot analysis (top). The blot was then stripped and reprobed with a GAPDH probe to control for the amounts of RNA loaded on the gel (bottom). Results are representative of five independent experiments.

Figure 6. Primer extension analysis of total RNA. 3 d after transfection, 3 μg of total RNA (1/50 of total RNA prepared from one 10-cm dish) was subjected to primer extension analysis as described. Primer extension products were electrophoresed on an 8% polyacrylamide-urea gel and autoradiography using an intensifying screen with exposure time of 24 h at −80°C was performed. A sequencing reaction using the same primer on the adw strain is shown on the left (lanes marked A,G,C,T). Lanes are as indicated. Results are representative of four independent experiments.
pregenomic RNA appeared to be the most abundant transcript (Fig. 6). The construct containing MT5/6 displayed an approximately twofold increase (1.96 ± 0.45, from four experiments) of the pregenomic RNA transcript compared to the wild-type strains or the constructs with individual mutation 5 or 6 (a typical experiment is shown in Fig. 6). The level of precore RNA transcripts was not influenced by MT5/6, compared with the wild-type. To confirm the minor effect of MT5/6 on core promoter activity, we inserted the wild-type and mutant (MT5/6) core promoters into the promotorless, luciferase gene plasmid pXp2. These constructs were transfected into HepG2 cells and luciferase expression was analyzed. Fig. 7 demonstrated that the core promoter mutation MT5/6 led to a 2.3-fold increase in the expression of the reporter gene.

In addition to regulating pregenomic RNA transcription, the HBV core promoter also directs transcription of precore RNAs. The precore RNAs are translated into the precore protein, which is posttranslationally processed and secreted as hepatitis B virus e antigen (HBeAg). As shown, primer extension analysis of total RNA did not reveal any significant difference in precore RNA transcription. To extend this observation, we studied the effect of MT5/6 on HBeAg synthesis.

Table I. HBV Protein Secretion in Medium of Cells Transfected with HBV Constructs

<table>
<thead>
<tr>
<th>Construct</th>
<th>hGH (ng/ml)</th>
<th>HBsAg (S/N)</th>
<th>HBeAg (S/N)</th>
</tr>
</thead>
<tbody>
<tr>
<td>adw</td>
<td>6.0</td>
<td>27.7</td>
<td>9.2</td>
</tr>
<tr>
<td>ayw</td>
<td>6.2</td>
<td>23.9</td>
<td>10.0</td>
</tr>
<tr>
<td>MT5/6</td>
<td>5.9</td>
<td>22.6</td>
<td>11.2</td>
</tr>
<tr>
<td>MT5</td>
<td>7.1</td>
<td>24.1</td>
<td>9.0</td>
</tr>
<tr>
<td>MT6</td>
<td>6.5</td>
<td>25.0</td>
<td>9.6</td>
</tr>
<tr>
<td>FH</td>
<td>5.0</td>
<td>15.6</td>
<td>1.3</td>
</tr>
</tbody>
</table>

Huh-7 cells were maintained and transfected with various constructs. Transfection efficiency was monitored by cotransfection with pTKGH and subsequent analysis of secreted hGH by radioimmunoassay. On day 3, culture media were harvested for detection of hGH, HBsAg, and HBeAg. Average values from hGH, HBsAg, and HBeAg radioimmunoassays are shown (for HBsAg and HBeAg a signal to noise ratio [S/N] of > 2 is considered positive). Results are representative of four independent experiments.

After transfection of HuH-7 cells with various wild-type and mutant constructs, secreted HBeAg and hGH (as control) were analyzed in the medium of the transfected cells. As shown in Table I, the identified core promoter mutations had no effect on the HBeAg synthesis. In contrast, the construct containing the fulminant strain led to the well-described absence of HBeAg synthesis (Table I). HBsAg production was not affected to any significant extent by any of the mutations (Table I).

MT5/6-induced enhanced replication is independent of HBX. Since the core promoter overlaps with the HBX gene, MT5/6 also affects the HBX gene. MT5/6 in the HBX ORF resulted in two amino acid substitutions in the HBX protein; valine is replaced by isoleucine in position 132 and phenylalanine is changed to tyrosine in position 133. Since the biological function of HBX is not completely understood, it is difficult to assess the true effect of MT5/6 on HBX function. The only established function of HBX is its ability to transactivate transcription of a variety of cellular and viral genes (20–22). To examine the effect of MT5/6 on HBX transactivation, wild-type

Figure 7. Effects of MT5/6 on core promoter activity. Promotor constructs containing no (negative control), adw core promotor, MT5/6 core promotor, or RSV promotor (positive control) driving the luciferase gene were transfected into HepG2 cells. 48 h after transfection, luciferase activities were measured as described. Results are representative of four independent experiments.
and mutant HBX constructs were compared in their ability to transactivate the luciferase reporter gene. After cotransfection of the HBX expression and reporter constructs into HepG2 cells, no appreciable difference in transactivation was detected between the wild type and mutants (Fig. 8).

To investigate whether the amino acid changes of the HBX protein are important for MT5/6-induced enhanced replication, MT5/6 was introduced into the second copy of the HBX coding sequence at the 3' end of the terminal-redundant construct adwR9 (Fig. 2). In this position MT5/6 did not show the phenotype of enhanced viral replication (data not shown). To further elucidate the role of mutated HBXs in MT5/6-induced enhanced replication, the HBX open reading frame was inactivated in the replication-competent construct containing MT5/6 at its 5' end (Fig. 9 A). HBX contains three conserved, in-frame start codons, each of which has been proposed to encode functionally related gene products (23, 24). Since the replication-competent R9 construct begins at nt 1417, it contains only the two translational initiation sites for the middle (p8) and small (p6.6) HBX at its 5' end. Inactivation of these reading frames was obtained by introduction of a stop codon and a mutated start codon for p8 (HBX.2) and p6.6 (HBX.3) as shown in Fig. 9 A. Analysis of replicative intermediates of these constructs transfected into HuH-7 cells demonstrated that MT5/6-induced enhanced replication was independent of HBX inactivation (Fig. 9 B).

MT5/6 increases viral encapsidation independent of pregenomic RNA transcription level. The following experiments were designed to further characterize the mechanism by which the mutation(s) affect HBV encapsidation. Encapsidation of the HBV pregenomic RNA is thought to require the e encapsidation signal on the pregenomic RNA, the core protein, and the polymerase protein (5, 6, 19). Since MT5/6 appears to be outside these three elements, it likely affects viral encapsidation in a novel manner. It is not known, whether other components, e.g., other viral or cellular factors, are involved in the encapsidation process. By acting in cis and affecting the core promoter function with a twofold increase in pregenomic RNA transcription level, MT5/6 could cause a modest increase in viral encapsidation. However, this effect on pregenomic RNA transcription (twofold) was much smaller than the observed effect on encapsidation (> 10-fold) of pregenomic RNA by MT5/6. Therefore, mechanisms other than increased level of transcription of pregenomic RNA must be present. A possible effect of MT5/6 on additional yet unidentified encapsidation sequences could be the reason for more efficient encapsidation. However, this possibility is highly unlikely because mutations introduced only in the 5' terminal redundant region of the R9 construct should not be present in the pregenomic RNA transcript (see Fig. 2).

To address this issue further, we applied a previously described encapsidation system (14). In this system a plasmid expressing a chimeric RNA (HBVe-lacZ) bearing the HBV e encapsidation sequence fused to lacZ is cotransfected with a plasmid expressing core and polymerase (pCMV-CP). The chimeric RNA was shown to encapsidate into core particles (14). This system would permit us to discern a direct cis effect of MT5/6 on pregenomic RNA transcription and encapsidation, or a transcript level-independent trans effect on encapsidation induced by MT5/6. To study the cis effect, we cotransfected the core promoter constructs pCPadwlacZ and pCPMT5/6lacZ together with pCMV-CP. MT5/6 resulted in increased encapsidation of HBVe-lacZ RNA by two to three times (2.4±0.5, from four experiments; Fig. 10 C, lanes 7 and 8). The enhanced encapsidation was due to increased transcription of the chimeric RNA (Fig. 10 C, lanes 1 and 2), which is consistent with our previous findings. Next, we provided the mutation(s) in trans by cotransfecting the wild-type construct pCPadwlacZ together with either adwR9 or MT5/6R9. In this experiment, MT5/6 led to a dramatic increase of encapsidation of HBVe-lacZ RNA (Fig. 10 C, lanes 9 and 10) without affecting the transcription of HBVe-lacZ RNA (Fig. 10 C, lanes 3 and 4). This blot was overexposed to reveal the weak signals of lanes 7 and 8. In a lighter exposure, the significant differences between lanes 9 and 10 as well as between 11 and 12 were much
more evident. Densitometric quantification revealed a >10-fold difference (10.9±2.1, from four experiments) between the wild-type and mutant signal intensities. A similar effect was also evident when a heterologous CMV promoter construct pE-lacZ was used instead of pCPadwLacZ (Fig. 10 C, lanes 11 and 12). In the pE-lacZ construct, the core promoter was replaced with the CMV promoter to drive the chimeric RNA. This experiment further supports the notion, that a major effect of MT5/6 on encapsidation is independent of the level of transcription.

**Discussion**

In our previous study, we demonstrated an HBV strain associated with an outbreak of fulminant hepatitis B exhibiting the phenotype of enhanced replication in tissue culture. We demonstrated that the increased level of core-associated HBV pregenomic RNA and replicative intermediates was associated with enhanced virion production in the medium (10). In this study, we defined the mutations of the FH strain responsible for the enhanced replication to the overlapping region of core promoter and distal HBX gene. The combined mutations of C to T at nucleotide position 1768 and T to A at nucleotide position 1770 were sufficient to induce the enhanced replication phenotype observed with the FH strain (Fig. 3). Similar to the FH strain, these mutations caused enhanced replication predominantly by an increased encapsidation of pregenomic RNA (Fig. 4). These two mutations were present in all the clones isolated from the five patients with fulminant hepatitis as well as the source patient who transmitted the virus (9). This observation indicated that these two mutations were not a result of PCR or cloning artifacts. Although other mutations in this viral strain may have other phenotypic effects, our data suggest that these two mutations resulting in enhanced replication are likely a unique feature of this naturally occurring HBV mutant.

Previous studies suggested that HBV replication is higher in postconfluent cells (15). It is possible that MT5/6 relieves the virus of this dependence to result in an overall higher replication. We have analyzed viral replication and observed enhanced replication of the mutant strain at various time points in culture. Thus far, mutations leading to enhanced replication have been described only in avian hepadnavirus. In duck hepatitis B virus, an amino acid change in the large envelope protein has been shown to enhance viral replication by increasing the cellular pool of ccc DNA (25).

Enhanced replication of the FH strain was the result of enhanced encapsidation of pregenomic RNA. Enhanced encapsidation was caused by two mechanisms: First, MT5/6 resulted in a two- to threefold increase in pregenomic RNA transcription, leading to a similar increase in viral encapsidation (Fig. 10). Second, MT5/6 led to a transcript-independent increase in encapsidation efficiency (>10-fold) by a yet unidentified mechanism (Fig. 10).

Increased transcription of pregenomic RNA was the result of altered core promoter function induced by the mutations. The core promoter regulates the transcription of the prege-
nomic RNA which is translated into the core protein and serves as the template for reverse transcription during viral replication (1). Primer extension analysis and Northern blot analyses of total RNA showed an approximately twofold increase in transcription of the pregenomic RNA in MT5/6 as compared with the wild-type adw strain (Figs. 5 and 6). Using a luciferase reporter, we observed a similar effect (2.3-fold increase) of the mutations on core promotor activities (Fig. 7). In addition to directing pregenomic RNA transcription, the core promotor regulates transcription of the precore RNAs which are translated into HBeAg (26, 27). Primer extension analysis of total RNA and analysis of HBeAg synthesis demonstrated that MT5/6 had no significant effect on transcription and expression of the precore gene. Recently, it was suggested that the precore expression product (HBeAg precursor protein) can inhibit HBV DNA synthesis by interfering with viral encapsidation (28). Since precore gene expression was not affected by MT5/6, the effect of MT5/6 on viral encapsidation via this mechanism seems to be unlikely.

In addition to increased transcription of pregenomic RNA, encapsidation was enhanced by a mechanism apparently independent of the level of pregenomic RNA transcription (Fig. 10). Encapsidation of pregenomic RNA has been shown to require the ϵ encapsidation signal present in the pregenomic RNA, the core, and the polymerase (5, 6, 14, 19). Our experiments suggest that the sequence in the core promotor-HBX gene may have a yet unidentified role in the regulation of the encapsidation process. Several mechanistic pathways are possible to explain the observed transcript-independent effect of MT5/6 on viral encapsidation.

First, encapsidation may be increased by a viral trans-acting factor. Such a trans-acting factor could be the HBX protein. Various forms of HBX protein, based on translation initiation at three different AUG start codons within the HBX open reading frame, have been postulated to exist (23, 24). The biological importance of the HBX protein(s) in viral life cycle remains largely unknown. Full-length HBX has been shown to be dispensable for HBV replication in tissue culture (29) but necessary for establishment of woodchuck hepatitis virus infection in vivo (30, 31). In transgenic mice, high level viral replication could be achieved only with a terminally redundant HBV transgene containing a duplicate HBX gene at its 5′ end (32). HBV transgenes without the HBX coding region in the 5′ end yielded no or low level HBV replication in mice (32). Although these transgenes contain an additional copy of the entire HBX gene at their 3′ end, this copy is apparently not sufficient to direct high level viral replication (32). In our system, MT5/6 placed in the 3′ terminal redundant region of the R9 construct had no effect on viral replication or encapsidation. Inactivation of HBX translation by oligonucleotide-directed mutagenesis did not abolish the MT5/6-induced enhanced replication (Fig. 9). In addition, introduction of MT5/6 in a HBX expression construct did not alter the transactivation of a luciferase reporter gene (Fig. 8). These findings essentially exclude an important role for the HBX gene product in mediating MT5/6-induced enhanced replication. Alternatively, transcripts or proteins of an antisense open reading frame (ORF) or spliced sequences such as those proposed earlier (30, 33) may mediate a trans effect on viral encapsidation. However, inactivation of an identified antisense open reading frame by inserting a frame-shift mutation did not abolish MT5/6 induced enhanced replication (data not shown).

Second, enhanced encapsidation could be the result of a cotranscriptional effect on core and polymerase expression induced by MT5/6. Such a cotranscriptional effect could result in a more efficient core or polymerase expression leading to an increased assembly of the encapsidation components. Translation of the RT-Pol appeared to occur through a novel ribosomal scanning mechanism (34, 35), probably depending on dynamic interaction of cellular initiation and elongation factors bound to the pregenomic transcript. Since the RT-Pol is probably the limiting factor for encapsidation, it is conceivable that MT5/6 alters the binding complex of these cellular factors during transcription leading to increased RT-Pol translation. MT5/6 may also act post- or cotranscriptionally by interacting with cellular factors required for RT-Pol activity and the subsequent encapsidation process. A cellular factor, Hsp90, has been recently identified to interact with the replication complex (36). Further experiments are under way to identify possible viral and/or cellular elements involved in MT5/6-induced enhanced encapsidation.

Regardless of the mechanism, our study demonstrates an important role for the core promotor sequence in HBV encapsidation and replication. Our current data, however, do not distinguish between whether MT5/6 exerts a positive or negative effect on this yet unknown element which could have either an activating or repressing role in encapsidation and replication. Further studies are necessary to clarify whether this finding is unique to this FH strain or it is important for HBV replication in general. Mutations in the core promotor region have been detected in patients with fulminant hepatitis B in Japan (37) and Europe (38) but less frequently in the U.S. (39). Our identified C-to-T mutation at nucleotide position 1768 and T-to-A mutation at nucleotide position 1770 were present in one of the Japanese strains (37). Core promotor mutations have also been described in HBV carriers and patients with chronic hepatitis (40). These studies led to the conclusion that core promotor mutations may influence the biological behavior of the virus, leading to a different clinical course. The authors of these studies speculated that transcription of the precore mRNA might be decreased as a result of these mutations leading to diminished HBeAg synthesis. Furthermore, they suggested that both the precore stop codon and core promotor mutation(s) are functionally equivalent resulting in a common phenotype of HBeAg deficiency. However, no functional studies have been performed to show that these mutations indeed have an effect on viral protein expression. Our study showed that HBeAg production was not affected by the specific core promotor mutations 5 and 6 (Table 1). Functional studies of the core promotor sequences of other viral strains associated with fulminant hepatitis are necessary to elucidate the effect of core promotor mutations on viral replication. Furthermore, we have no direct evidence that the enhanced replication of the FH strain contributes to the development of fulminant hepatitis. Studies in an animal model, such as in woodchuck, would be crucial in demonstrating the biological significance of these mutations in the pathogenesis of HBV infection.

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