X-deficient woodchuck hepatitis virus mutants behave like attenuated viruses and induce protective immunity in vivo

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The X protein (HBX) of the hepatitis B virus (HBV) has been shown to be important for the establishment of HBV infection in vivo. Our previous studies suggested that interaction of HBX with the proteasome complex may underlie the pleiotropic functions of HBX. In this study, we generated a series of woodchuck hepatitis virus (WHV) X mutants, including mutants of the domain interacting with the proteasome, and studied their infectivity in woodchucks. Many of the mutants were defective in transactivation but none of them were completely replication defective in vitro. In vivo, all the wild-type and some X mutant–transfected animals demonstrated evidence of infection with anti-WHc and/or anti-WHs seroconversion. Most of the wild-type– and X mutant–transfected animals had transient viremia. Some animals were later challenged with infectious WHV. Animals inoculated with X mutants, including those with no serologic evidence of infection, were protected from the challenge, suggesting previous infection with resulting protective immunity. Our study demonstrates that the previously described functional domains of HBX are biologically important and the X-defective mutants, possibly as attenuated viruses, are not completely replication defective in vivo.


Introduction

Human hepatitis B virus (HBV) is a member of the Hepadnaviridae family, which includes the hepatitis viruses of the woodchuck, ground squirrel, tree squirrel, Pekin duck, and heron. HBV has a unique fourth open reading frame, termed the hepatitis B virus X (HBX) gene. The HBX gene is well conserved among the mammalian hepadnaviruses and codes for a 16.5-kDa protein. The protein can activate the transcription of a variety of viral and cellular genes (1, 2) and induce liver cancer in certain transgenic mouse models (3). Since HBX does not bind to DNA directly, its activity is thought to be mediated by protein-protein interactions. HBX has been shown to enhance transcription through AP-1 and AP-2 (4, 5) and to activate various signal transduction pathways (6, 7). Several recent studies have also identified possible cellular targets of HBX, including members of the CREB/ATF family (8), the TATA-binding protein (9), RNA polymerase subunit RPB5 (10), the ultraviolet-damaged (UV-damaged) DNA-binding protein (11), and the replicative senescence protein p55sen (12). HBX has also been shown to interact with p53 and inhibit its function (13, 14).

Using the Saccharomyces cerevisiae two-hybrid system and a modified yeast two-hybrid dissociator system, we have recently identified two proteasome subunits — PSMA7 (an α proteasome subunit) and PSMC1 (an ATPase-like member of the 19S regulatory factors) — as putative cellular targets of HBX (15, 16). We demonstrated that this interaction may be functionally important in the pleiotropic effect of HBX (17). Because woodchuck hepatitis virus (WHV), like HBV, encodes an X protein (WHVX) with transactivation function and WHVX also interacts with the two proteasome subunits (15), infection of woodchucks would be a valuable model for studying the WHVX function in vivo. In this study, we created a series of site-directed WHVX mutants, including mutants of the domain interacting with the proteasome subunits, and studied the in vivo behaviors of these mutants in woodchucks.

Methods

Plasmid construction. A 1.2× full-length genome of an infectious WHV strain (pWHV8) was used in this study (18). Various WHVX mutants (Figure 1) were generated by PCR mutagenesis (QuickChange Site-Directed Mutagenesis Kit; Stratagene, La Jolla, California, USA) and confirmed by sequencing: WHV8Xlg– with ATG→TTG mutation at the first translation initiation site of WHVX, WHV8Xmd–, and WHV8Xsm– with ATG→GTG mutations at the second and third in-frame translation initiation sites, respectively; WHV8G67S with Gly→Ser at amino acid (aa) 67, WHV8P68A with Pro→Ala at aa 68, WHV8G132V with Gly→Val at aa 132, and WHV8H135D with His→Asp at aa 135. The WHV8XRN–
was created by introducing a new in-frame TTG codon to ATG at aa 55 in the X gene. These X mutations do not cause any amino acid changes in the polymerase gene overlapping the X gene. A replication-defective WHV control, WHV8Pmt, was created by mutating the Asp–Asp→Ala–Ala at the YMDD motif of the polymerase gene. The introduction of these two mutations has been shown previously to inactivate the Pol gene (19, 20).

**DNA transfection and transactivation assay.** Luciferase (Luc) reporter construct RSV-Luc consisting of Rous sarcoma virus promoter driving the luciferase gene was used for WHVX transactivation assays in the HepG2 cells as described previously (15, 16). The human hepatoma HepG2 or HuH7 cells were grown in DMEM (Life Technologies Inc., Rockville, Maryland, USA) containing 10% FBS in a humidified incubator (5% CO2). Transient transfection of HepG2 cells in a 35-mm well was carried out using the DNA transfection kit (5 prime→3 prime Inc., Boulder, Colorado, USA). Luciferase assay was performed with a Monolight Luminometer (Analytical Luminescence Laboratory, San Diego, California, USA). All transfections were controlled for transfection efficiency by cotransfecting a pTK-GH plasmid as described previously (15).

**Viral nucleic acid analyses.** HuH7 or HepG2 cells were transiently transfected with WHV8 clones described above. Three or four days after transfection, the cells were harvested for viral RNA and DNA analyses. RNA was prepared by the guanidinium isothiocyanate-acid-phenol method (ULTRASPEC RNA isolation kit; BIOTECX Laboratories, Houston, Texas, USA), analyzed by 1% formaldehyde agarose gel electrophoresis, and hybridized with [32P]-labeled WHV-specific probe. Viral replicative intermediates associated with intracellular core particles were isolated by ultracentrifugation of cell lysate through a 30% sucrose cushion and analyzed by Southern blot hybridization as described (21).

**Intrahepatic inoculation of woodchucks with WHV DNAs and challenge of woodchucks with serum-derived WHV7.** All the experimental woodchucks were offspring of WHV-free parents that were born and raised in the animal facility of the College of Veterinary Medicine at Cornell University. For intrahepatic inoculation of woodchucks, various WHV8 constructs were purified by standard protocols (Plasmid Maxi Kit; QIAGEN Inc., Valencia, California, USA). Three animals were transfected with each mutant, and six animals were inoculated with wild-type WHV8 DNA as positive controls. To avoid cross-contamination, inoculations with wild-type and mutant constructs were performed on different days, and the animals were kept at different containment rooms. Each woodchuck received a dose of 50 µg plasmid DNA in a volume of 0.5 ml of PBS. Starting 2 weeks after transfection, serum samples were collected biweekly for a total of 24 weeks. Liver biopsies were performed every 4 weeks. At week 24 after transfection, selective woodchucks were challenged with 107 infectious dose of serum-derived WHV7. Serum samples were collected biweekly until 12 weeks after challenge.

**Serological and nucleic acid assays.** For markers of WHV infection, serum samples collected serially were tested for the presence of WHsAg and Ab’s against WHV core (anti-WHC) or sAg (anti-WHS) using solid-phase radioimmunoassay or ELISA. WHV DNA was assayed routinely by the slot blot hybridization method. For the detection of low viremia of the transfected woodchucks, WHV virions from woodchuck serum were concentrated by ultracentrifuge through a 30% sucrose cushion and treated with micrococcal nuclease to eliminate contaminating free nucleic acids. WHV DNAs were purified by proteinase K digestion and phenol extraction as described previously (21). The virion DNAs were amplified with WHVX-specific primers by PCR with Herculase Enhanced DNA polymerase (Strategene). Several known negative serum samples were included in each assay, and if any of them became positive, the results were discarded and the assay repeated.

**Figure 1**
Schematic diagram of WHVX mutants. The WHVX mutants are displayed in reference to the genomic map of WHV8. The shadow regions represent the two important structural domains of WHVX. The mutations and their approximate positions are shown. The nucleotide sequences are shown in the middle, the amino acids of the mutations are shown at the bottom for the X gene and at the top for the overlapping Pol gene (P gene).
Amplification, cloning, and sequencing of WHV DNA from woodchuck samples. To avoid minor contamination of mutant WHV constructs with wild-type DNAs, the mutant DNAs were produced using disposable glassware at a different laboratory and on different days from that of the wild-type construct. The mutant constructs that were used for inoculation were further analyzed using the amplification refractory mutational detection system, which is based on the absolute inability of an oligonucleotide with mismatched 3′ residue to function as PCR primer under appropriate conditions (22, 23). In brief, an oligonucleotide primer of the wild-type sequence was designed to specifically amplify the wild-type sequence at the particular mutational site. For the Xlg− mutation, the wild-type primer Xwt.F, 5′ AAGCGTACGCCTTTACA 3′ (nt 1486 to 1503), together with the downstream primer X.R, was designed for the specific detection of wild-type sequence under the PCR condition of 94°C for 30 seconds, 60°C for 30 seconds, and 72°C for 1 minute using the hot-start Pfx DNA polymerase (Life Technology Inc.).

The virion DNAs were purified as described above and amplified with two pairs of WHV primers: the first pair (PreX.F primer, nt 1392 to 1410: 5′ TATTGCGCGGGGAACTC 3′; X.R primer, nt 1935 to 1914: 5′ TACATGGT- TACAGAAGTCGCAT 3′) amplifies the N terminus of WHVX, whereas the second pair (S.F primer, nt 1748 to 1767: 5′ ATTCCACCGTGAACTTTGTC 3′; C.R primer, nt 2346 to 2327: 5′ GTGAGACATGACAAATGAAA3′) amplifies the C terminus. To distinguish the virion DNA and the cytomegalovirus promoter–driven (CMV-driven) plasmid DNA, a CMV-specific sense primer (CMV.F primer, 5′ AAATGGGCGGTAGGCGTGTA3′) and the X.R primer were used (see Figure 5). The amplified fragments were cloned into plasmid vector pCR4Blunt-TOPO (Invitrogen Corp., Carlsbad, California, USA) and sequenced.

Results
Construction of WHV mutants. HBX has been proposed to contain three in-frame translation initiation sites that may result in multiple polypeptides with different functions (24). WHVX, like HBX, also contains three alternative translation initiation sites. We mutated the three sites separately, resulting in three WHVX mutants, named Xlg−, Xmd−, and Xsm− (Figure 1). Both HBV and WHV X genes contain two important structural and functional domains, with the second domain obligatory for interaction with the proteasome (15, 16). We created two first-domain mutants, G67S and P68A, and two second-domain mutants, G132V and H135D, respectively (Figure 1). Both HBV and WHV X genes contain two important structural and functional domains, with the second domain obligatory for interaction with the proteasome (15, 16). We created two first-domain mutants, G67S and P68A, and two second-domain mutants, G132V and H135D, respectively (Figure 1). We have shown previously that these mutations do not affect the stability of HBX (15). The N terminus of the X gene has been suggested to contain a regulatory domain of about 50 aa (25). To study the function of this domain in vivo, we created a mutant, XRN−, by altering the first ATG to TTG and a TTG codon at aa 55 to ATG, which is in a proper Kozak context for translation initiation codon (26). This mutant was shown to direct the production of a smaller X by in vitro translation (not shown). A replication-defective WHV control, WHV8Pmt, was generated by mutating the Asp-Asp→Ala-Ala at the YMDD motif of the polymerase gene (19, 20).

Transactivation by WHVX mutants. To test the transactivation function of the WHVX mutants, HepG2 cells were cotransfected with the WHV expression constructs and the reporter construct RSV-Luc. The wild-

![Figure 2](image-url)

**Figure 2** Functional characterization of WHVX mutants in vitro. (a) Transactivation of WHVX mutants. The reporter plasmid RSV-Luc and WHVX mutants were cotransfected at a ratio of 1:5 with a total DNA of 0.6 µg per well in a six-well plate. Luciferase activities were assayed 3 days later. Data shown are the mean values of triplicates. (b) Transcription of WHVX mutants. Three days after transfection, HuH7 cells were harvested, and total RNAs were isolated and subjected to Northern blot analysis. The ribosomal RNAs (bottom) were shown for RNA-loading control (probing for β-actin showed similar results). (c) Replication of WHVX mutants. Three days after transfection, WHVX-transfected HuH7 cells were harvested, and the core particle–associated WHV DNA were isolated and subjected to Southern blot analysis. Lanes are as indicated. SS, single-stranded WHV DNA. All the results are representative of at least three independent experiments.
type WHV exerted a 20-fold transactivation of the reporter gene, the WHV8Pmt functioned in a manner similar to the wild-type, and the md–, sm–, P68A, and RN– constructs transactivated at about 30–60% of the wild-type level. All other mutants had little or no transactivation function (Figure 2a).

Transcription and replication of WHVX mutants. To evaluate the WHVX mutants at the transcription and replication levels, viral nucleic acids were analyzed in transfected HuH7 cells. Compared with the wild-type WHV8, the viral transcripts of Xlg–, G132V, H135D, and RN– were reduced about 50%. All other mutants appeared to have WHV RNA levels similar to wild-type (Figure 2b). For the analyses of viral replication, viral DNA replicative intermediates associated with intracellular core particles were isolated and analyzed by Southern blot hybridization (Figure 2c). Compared with the wild-type WHV8, the RN– mutant replicated similarly, but all other mutants exhibited variable reductions in the level of replicative intermediates, ranging from about 10% (Xlg–) to 60% (md– and sm–). As expected, the WHV8Pmt mutant showed no replication (Figure 2c).

Infectivity of WHVX mutants in woodchucks. To study the infectivity of WHVX mutants in woodchucks, WHV-free adult woodchucks were transfected with the WHV8X mutants and tested for WHV serological markers and serum WHV DNA posttransfection (Figure 3). For the analyses of viral replication, viral DNA replicative intermediates associated with intracellular core particles were isolated and analyzed by Southern blot hybridization (Figure 2c). Compared with the wild-type WHV8, the RN– mutant replicated similarly, but all other mutants exhibited variable reductions in the level of replicative intermediates, ranging from about 10% (Xlg–) to 60% (md– and sm–). As expected, the WHV8Pmt mutant showed no replication (Figure 2c).

Figure 3
Infectivity of WHVX mutants in woodchucks. WHV-free adult woodchucks were transfected with WHV8X mutants, and serological markers were tested biweekly after transfection. Open bars depict samples with no serological evidence of WHV infection, while filled boxes depict serological evidence of WHV infection with anti-WHc and/or anti-WHs seropositivity. At week 24 after inoculation, selected woodchucks were challenged with 10^7 WID_{50} of WHV7 and followed for evidence of protection. The details of the challenge experiments are described in Figure 6, and the results are summarized here. +, protection; −, no protection; ND, not done.

Figure 4
PCR analyses of WHV DNAs. (a) Standard PCRs were performed with three pairs of primers: Pre-X.F and X.R pair (NX pair) to amplify N terminus of X with expected PCR product of 543 bp’s; X.F and C.R pair (CX pair) to selectively amplify C terminus of X of viral DNA (expected PCR product: 597 bp’s); CMV.F and C.R pair (CMVX pair) to amplify only cloned DNA (expected PCR product: 504 bp’s). The positions of these primers in the viral DNA and cloned DNA are shown. (b) Amplification refractory mutation detection system for wild-type sequence in X mutant inoculum DNA. The position of the wild-type–specific primer Xwt.F is shown in a. Wild-type and Xlg– constructs were mixed at ratios of 1:10 to 1:10,000. One nanogram of each mixture and the mutant inoculum was amplified for 30 cycles as described in Methods. (c) PCR-Southern analyses of viral and cloned WHV DNAs with three pairs of primers: lanes 1 and 4 with NX pair; lanes 2 and 5 with CX pair; lanes 3 and 6 with CMVX pair. The viral DNA was extracted from WHV-containing serum of known titer. About 1 pg of viral or cloned DNA was used for the analyses.
ure 3). All the wild-type–transfected animals demonstrated seroconversion and five of six of the animals had transient low-level viremia. Surprisingly, two of four Xlg–transfected animals exhibited anti-WHc and/or anti-WHs seroconversion. A fraction of other X mutant–transfected animals also displayed serologic evidence of infection. Most of the X mutant animals also had transient viremia regardless of serologic status. Most surprisingly, one of the three Pmt-transfected woodchucks also developed anti-WHc. However, none of the Pmt-transfected animals had demonstrable viremia (Table 1). Therefore, the presence of anti-WHc in this animal is probably a result of DNA immunization.

Analyses of WHV sequences. To eliminate the potential contamination of wild-type with mutant WHV construct during the DNA preparation stage, a wild-type–specific primer was designed for the amplification refractory mutation detection system (22, 23). The assay showed that no contamination of wild-type could be detected at a ratio of 1:10,000 (Figure 4b). Because 50 µg of WHV plasmid DNA was injected into the animals, it is possible that any detected WHV DNA in the serum could be a result of this residual DNA. Therefore, we developed a strategy to distinguish between circulating virion DNA and cloned plasmid DNA (Figure 4, a and c). First, we purified the viral particles by ultracentrifugation in a sucrose gradient and then treated the particles with micrococcal nuclease to eliminate all residual contaminating nucleic acids. Control experiments with spiked cloned WHV DNA in the serum showed at least a 6-log depletion of the plasmid DNA using this method (not shown). Second, we developed a selective PCR method to distinguish the two forms of WHV DNAs with three pairs of primers: PreX.F and X.R pair to amplify N terminus of X (NX) in both viral and cloned DNAs, X.F and C.R pair to amplify the C terminus of X (CX) virion DNA only, and CMV.F and C.R pair to amplify cloned DNA

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*aMutant sequence. *Mixture of mutant and wild-type sequences, confirmed by both direct sequencing of PCR products and cloning followed by sequencing of at least 20 clones. All the other detected sequences are wild-type. W.C., woodchuck.
immunologic priming. Two animals (WHV8 6016 and
G132V 6061, and H135D 6066) were protected, sug-
mitive serology were protected from the challenge: wild-
ture 3 and 6). Given the small amount of DNA injected,
none of them were protected from the challenge (Fig-
which was likely a result of DNA immunization, but
A WHV mutant with
antibodies in Pmt 6137. Detailed serological profiles of some
of the challenged animals are summarized in Figure 6.

Discussion
Because of the essential role of the X gene in the life

cycle of mammalian hepadnaviruses in vivo, character-
ization of the structural and functional properties of X
protein in an infectious animal model is crucial. While
the avian hepadnavirus may encode an X-equivalent
gene (29), the X gene of WHV has been studied in the
woodchuck, which probably represents the best small
animal model to address the issue. We have demon-
strated previously the structural and functional inter-
actions between HBX and the proteasome complex and
characterized the domain of HBX that is important for
this interaction with a well-defined structural and
functional correlation (15–17). In this study, we intro-
duced various single amino acid mutations into the
domains of HBX that have been shown to be function-
ally important for X in vitro, including the domain
interacting with the proteasome. In adult woodchucks,
these X mutants appeared to behave like attenuated
viruses with low-level replication and anti-WHV sera.

Challenge with infectious WHV. At 24 weeks after trans-
fection, selective woodchucks were challenged with a
10^6 infectious dose of serum-derived WHV7. The serum
samples were collected and tested for serological evi-
dence of protection against WHV infection. The data
are summarized in Figure 3. All the animals with posi-
tive serology were protected from the challenge: wild-
type animals 6032, 6044, and 6016; Xlg - 6059 and 6055;
G67S 6019; and H135D 6010. Surprisingly, even wood-
chucks with no serologic evidence of infection (Xlg -
6046 and 5003, XRN - 6041 and 6070, P68A 6045,
G132V 6061, and H135D 6066) were protected, sug-
gest ing establishment of low-level infection with
immunologic priming. Two animals (WHV8 6016 and
Xlg - 6059) had weakly positive WHV DNA as deter-
mined by PCR in serum at postchallenge week 2 only,
which is likely a result of carryover from the inoculating
dose. Because 10^7 WID50 of WHV (about 10^8 genomes)
was inoculated, a small amount of WHV might remain
2 weeks later. If one assumes the half-life of free virion
in circulation is 1–2 days (27, 28), approximately 10^7
genomes/ml should remain after 2 weeks (~10 half-
lives), which is at the detection limit of our assay. We
also challenged four naïve animals (animals 2962, 2956,
4039, and 4094), and three Pmt-inoculated animals
(animals 6128, 6137, and 6167). Both naïves and Pmt
animals were not protected against the challenge and
developed transient viremia with positive WHsAg and
WHV DNA in serum, despite the presence of anti-WHc
Ab’s in Pmt 6137. Detailed serological profiles of some
of the challenged animals are summarized in Figure 6.

Figure 5
Sequencing and cloning of PCR product. Serum samples from chal-

lenged animals were analyzed by both direct sequencing and cloning
followed by sequencing of the PCR products. Direct sequencing (a)
and cloning/sequencing (b) of the PCR products from P68A-trans-
fected animal 6017 at weeks 4, 6, and 8 after challenge are shown as
an example. The sequences were confirmed by three independent

serum extraction/PCR analyses.

a

b

Week 4

Week 8

Week 6

Direct sequencing

Cloning/sequencing

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WT

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and it is capable of directing a first-round synthesis of pregenomic RNA independent of the endogenous HBV promoter. This construct has also been used to test the effect of X mutations in woodchucks, and the study also failed to show any infectivity of the mutant (32). However, in the study of Zoulim et al., only one animal was tested for the X mutant, and in our study we tested four animals. Furthermore, the above two studies depended on serology for infectivity determination and did not look for low-level viremia by PCR. As shown in our study, several animals inoculated with WHVX mutants did not develop anti-WHV Ab's but had demonstrable viremia. Finally the challenge study, performed only in our study, further supported the interpretation that X-deficient mutant can replicate at low level in vivo.

Analyses of WHV sequences during the acute phase of infection revealed the reversion of X mutants to the wild-type sequence except in the animals inoculated with the Xmd− and Xsm− mutants. Although Xmd− and Xsm− mutants had diminished transactivation functions and replicated at a reduced efficiency in cell culture, they seemed to behave similarly to the wild-type in vivo. Therefore, the notion of alternative initiations of X gene may not be biologically relevant (24). However, there may be subtle phenotypic differences that could not be discerned in this study. In animals inoculated with other X mutants, the genotypic reversion to wild-type supports the interpretation that early replication of the X mutants did occur in the woodchucks; otherwise the wild-type revertant would not have a chance to emerge. The temporal sequence of the reversion of X mutants to wild-type occurred rather quickly during the first 2 weeks (Table 1). Most of the X mutant–inoculated animals had predominantly the wild-type sequence in circulation by week 2. Only the animals inoculated with X mutant, XP68A, demonstrated a mixture of mutant and wild-type sequences.

**Figure 6**
Challenge of the transfected woodchucks with WHV7. At 24 weeks after transfection, selective woodchucks were challenged with a 10^7 infectious dose of serum-derived WHV7. Serological markers were tested biweekly from 4 weeks before challenge (weeks –4 and –2) to 12 weeks after challenge. Time 0 with an arrow represents the time of challenge. The serologic profiles of some of the challenged animals are shown. Signal-to-noise (mean OD of three serum samples of the animal before intrahepatic inoculation) ratio (S/N) for each serologic assay is shown. S/N greater than 3 is considered positive. Serum WHV DNA was assayed as described in Methods, and duplicate assays were performed independently for each serum sample. Only samples positive for both assays were scored as positive. For sample with only one positive assay, a third was performed; if the third was positive, the sample was scored as positive.
sequences in the first few weeks, which was then replaced completely by the wild-type sequence in circulation (Figure 5 and Table 1). Although a minor population of mutant species might still exist, we could not detect them based on the sensitivity of our assay (<1 of 20).

Based on the calculation of the rate of replication in vivo and the mutational frequency of the error-prone reverse transcriptase, a high probability of the emergence of mutants could be predicted in vivo (28). At the peak of hepadnaviral infection, all possible single nucleotide changes could occur in one day. However, the emergence of a viral species is determined by many factors, one of which is the replication fitness of the viral species (28, 33). Because the wild-type replicates more efficiently than the X mutants, it is expected that the wild-type can quickly replace the mutants, especially during the early phase of infection, in which there is little competition for replication space (34). It is not clear why only the XP68A persisted longer than the other X mutants in vivo. This X mutation may confer a less deleterious effect on the in vivo replication of WHV, as reflected in its partially active transactivation function in vitro (Figure 2a). This possible explanation awaits further experimentation.

The X mutants, while replicate with an attenuated phenotype, are capable of priming the immune response to protect the animals from subsequent challenge. All the animals demonstrated an anamnestic response of anti-WHc with a X-null mutant. In our study, the Xlg– mutant behaved similarly to all the X mutants with a X-null mutant. In our study does not permit a phenotypic comparison from these two studies that several domains of X are functionally important in vivo, but the precise roles of these interacting factors with X and productive viral infection await further experimentation.

Our study suggests that X-deficient virus, as an attenuated virus, could be considered as a vaccine candidate because it is capable of inducing protective immune response in vivo. In addition, the intriguing possibility of HBX as a potential target of antiviral development could greatly expand the armamentarium of anti-HBV therapy, which, to date, only consists of interferon and lamivudine.

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