Effective inhibition of hepatitis B virus replication by small interfering RNAs expressed from human foamy virus vectors

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Abstract. RNA interference (RNAi) mediated by double-stranded small interfering RNA (siRNA) is a novel mechanism of sequence-specific, post-transcriptional gene silencing. There has been much research into the use of RNAi for the treatment of human diseases. Many viruses, including hepatitis B virus (HBV), are susceptible to inhibition by this mechanism. However, for RNAi to be efficacious therapeutically, effective RNAi targeting sequences and a suitable delivery system are required. In this study, we employed a polymerase chain reaction (PCR)-based siRNA expression strategy to rapidly screen for effective siRNA sequences. Two effective siRNAs sequences (designated as S2 and X1) which reduced the HBV RNA by >90% were identified. For delivering the siRNAs, they were cloned into a human foamy virus (HFV)-based vector to generate single siRNA expression vectors HFVU6-siS2, HFVU6-siX1 and a dual siRNA expression vector HFVU6-siSX. The results showed that these siRNA vectors effectively inhibited multiple HBV gene expression and viral DNA replication based on ELISA and quantitative PCR analysis. HFVU6-siSX which simultaneously expressed two siRNAs that targeted the S and X genes of HBV is the most potent inhibitor of HBV replication. In addition, the repression of HBV RNA and DNA was stable for up to 3 months post-transduction as determined by RT-PCR and Southern blotting. Collectively, the PCR-based siRNA expression strategy provides a rapid and easy approach for testing candidate anti-HBV siRNA sequences and for cloning selected siRNA expression cassettes into a vector. RNAi based on the HFV vector was able to achieve effective, long-term inhibition of HBV gene expression and viral DNA replication. The combination of the two techniques may provide a powerful tool in the treatment of viral infection.

Introduction

Hepatitis B virus (HBV) infection is one of the major threats to public health. It is estimated that, worldwide, approximately 2 billion people have been infected and about 350 million have chronic HBV infection even though effective vaccines are available (1-4). These carriers are at high risk for developing HBV-associated diseases, including chronic hepatitis, liver cirrhosis and primary hepatocellular carcinoma. Worldwide, more than 1 million carriers die from these diseases each year (1). To date, interferon and nucleoside analogue combinational treatment has had only limited success. Thus, new therapeutic approaches are urgently needed.

RNA interference (RNAi) is a natural process in eukaryotic cells by which double-stranded RNA initiates and directs sequence-specific, post-transcriptional silencing of homologous genes. It is believed to have evolved as a host defense mechanism directed at transposable elements and infecting viruses (5,6). Recently, RNAi has been applied to inhibit the replication of several pathogenic human viruses. HBV appears to be an attractive target of RNAi, as its pregenome (pg) functions as both mRNA and replication template in the viral life cycle, all of its transcripts have common 3' ends, and multiple viral RNAs can be inhibited by a single short interfering RNA (siRNA). Therefore, RNAi is well suited for developing novel treatments for HBV infection.

Successful application of RNAi in the treatment of HBV infection primarily depends on two important factors. One is to screen specific RNAi targeting sequences which could effectively knockdown HBV transcripts. Konishi et al and McCaffrey et al have demonstrated that siRNAs targeted to different sequences of the HBV genome varied in their overall inhibiting effect, and only a limited number of siRNAs were capable of inducing highly effective inhibition (7,8). In
our study, a PCR-based siRNA expression strategy was employed to rapidly screen for effective siRNA sequences. Briefly, the siRNA expression constructs containing the U6 promoter and short hairpin DNA were established by overlap PCR, and the PCR products were directly transfected into cells resulting in functional expression of siRNAs. The other factor is to develop an efficient intracellular delivery system for siRNAs or the vectors expressing the corresponding siRNAs. Methods relying on lipid-based transfection reagents to introduce synthesized siRNAs or vectors into cells are either inefficient or unsuitable for use in animals. Viral vectors are good candidates for transducing siRNAs, and here a novel HFV vector was utilized to construct siRNA expression vectors. The HFV vector offers the advantages of efficiently infecting the cells and producing stable gene expression in the infected cells. In this study, we report the effects of HFV vector-directed siRNAs on HBV gene expression and replication.

Materials and methods

Targeting sites of siRNA on HBV genome. From the complete genome of HBV (GenBank Accession no. U95551), nine regions were selected as the targeted sequences using siRNA target finder program of Ambion (http://www.ambion.com//techlib/misc/siRNA_finder.html). The selected sequences were subjected to global BLAST to ensure that they did not have significant sequence homology with human genome sequences. These sequences targeting the position of the core region of 230, 268, and 306; S region of 69, 302, and 609; and X region of 271, 283, and 306 were as follows (Fig. 1A): C1, 5'-AAGATCCAGCATCTAGAGACC-3'; C2, 5'-AAGTTCAAGGCAACTCTTGTGG-3'; C3, 5'-AAGAGAAACCGTTATAGAGTA-3'; S1, 5'-AAGAATCCTCACAATACCGAC-3'; S2, 5'-AAGGTATGTTGCCCGTTTGTC-3'; S3, 5'-AAGGTCTGTACAGCATCTTGAG-3'; X1, 5'-AAGGTCTTACATAAGAGGACT-3'; X2, 5'-AAGAGGACTCTTGACTCTCT-3'; and X3, 5'-AATGTCAACGACCGACCTTGA-3'. Based on the previous report, a random siRNA (5'-GCGCGCTTTGTAGGATTCG-3') was selected as negative control siRNA (NC) (7).

Preparation of PCR products containing U6 promoter and shDNA. Human U6 promoter was amplified using the plasmid pAVU6+27 (9) as a template with the pair of primers: PU6F, 5'-CATATCCGGAAGGTCGGGCAGGAAG-3'; and PU6R, 5'-GGTGTTTCGTCCTTTCCACAAGATATAT-3'. Primer PU6F contains the AccIII restriction site underlined in the sequence, which enables directional cloning of PCR products into viral vectors. Human U6 promoter followed by a 19-nt sense strand of siRNA, a 9-nt loop, a 19-nt anti-sense strand of siRNA, and a stretch of six deoxythymidines were amplified by two-step overlap PCR, as previously described (10,11). In the first round of PCR, the reverse primer (3' primer 1) consisting of (5') the 9-nt loop complementary sequence (5'-TCTCTTGA-3'), a 19-nt anti-sense sequence and a complementary sequence to the 11 nt at the 3'-end of the human U6 promoter (3') was used. The first PCR reaction mixture (1 μl) was reamplified in the second round of the PCR reaction using the same forward primer PU6F and a new reverse primer, 3' Primer 2, which contained (5') poly(A)19, a 19-nt sense strand of siRNA, and a complementary sequence to the 9-nt loop (3') (Fig. 1B). The PCR products (U6-shDNA) were co-transfected into HepG2 cells. The levels of HBV RNA and luciferase mRNA were quantified by real-time PCR. The reduction in the ratio of HBV RNA to luciferase mRNA indicates the siRNA activity. The values shown are the average of three independent experiments (means ± standard deviation).

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Cell culture and transfection. HepG2 cells were maintained in DMEM supplemented with 10% fetal bovine serum at 37°C. HepG2 cells were seeded at a density of 1.0x10⁵ cells per well into 24-well plates and grown to ~80% at the time of transfection. Co-transfection of 0.1 μg of luciferase expression plasmid pGL3-Control (Promega, WI, USA), 0.5 μg of HBV replication competent plasmid pHBV (12) and 0.4 μg of the PCR products was carried out with Lipofectamine 2000 (Invitrogen, MD, USA). Forty-eight hours post-transfection, cells were harvested and total RNA was isolated using an RNeasy kit (Qiagen) with DNase treatment.
Generation of siRNA expression HFV vectors and transduction. The construction of pGPSNI-GFP and helper pΔGP have been described previously (13-15). The pHFVU6-siS2 and pHFVU6-siX1 plasmids were constructed by replacing the SV40 promoter with the corresponding siRNA expression cassettes into the restriction enzyme sites of AccIII and SmaI on pGPSNI-GFP. To generate the dual siRNA expression plasmid pHFVU6-siSX, the AccIII site of plasmid pHFVU6-siX1 was blunted and the U6-shS2 fragment was cloned into this site (Fig. 2A). The control plasmid pHFVU6-siNC was constructed with the random siRNA sequence as described previously. All constructs were confirmed by sequencing.

A series of HFVU6 vectors were produced by co-transfecting 293T cells with vector plasmids and helper plasmid pΔGP. Supernatants were harvested 4 days post-transfection and purified by centrifugation at 5000 rpm for 20 min. The vectors were then concentrated by using the Amicon Ultra-15 centrifugal filters, 100 k (Millipore, MA, USA). The titers of HFVU6 vectors were determined on 293T cells. Four to five days after infection, cells were monitored and scored for GFP fluorescence under a fluorescence microscope (Olympus IX71). Transduction of 2.2.15 cells was carried out at a multiplicity of infection (MOI) of 30 as previously described (13).

ELISA assays and immunofluorescent staining. The levels of HBsAg and HBeAg in the media were determined using the AxSYM systems kit (Abbott, IL, USA) based on enzyme-linked immunosorbent assay (ELISA) (16). All assays were performed three times independently.

The HBx protein was detected by immunofluorescent staining. Briefly, cells were washed and then fixed with 3% paraformaldehyde in phosphate-buffered saline (PBS) and permeabilized with 0.5% Triton X-100 in PBS for 10 min. After blocking with 3% bovine serum albumin (BSA) in PBS, cells were incubated with a monoclonal HBV X-protein antibody (Chemicon, CA, USA), followed by Cy3-conjugated secondary antibody (Sigma, MO, USA). Microscopic images were obtained with an Olympus fluorescence microscope.

Quantitative PCR analysis. Real-time PCR was performed to quantify HBV viral genomic DNA or RNA by using the HBV fluorescence quantitative PCR diagnostic kit (PG Biotech, Shenzhen, P.R. China). To quantify luciferase mRNA, primers 5'-GCCGGGTTTTTCTTTGTGAC-3'; reverse, 5'-GTGCTCCAGGGAAGTCC-3', and probe FAM-5CTGAAGTCACATTCTCCACATG-3' were used. As a negative control for DNA contamination, PCR was also performed on the sample without conducting the RT step. Products were analyzed by electrophoresis on a 2% agarose gel.

Southern blot analysis. DNA was extracted with the QIAamp DNA mini kit (Qiagen), then digested and electrophoresed in 0.7% agarose gels and transferred to Hybond-N+ membranes for Southern blot analysis. Full-length HBV DNA fragment was labeled with 32P and used as the hybridization probe. Hybridization signals were analyzed using Phosphor-Imager 830 (Molecular Dynamics, Sunnyvale, CA).

Results

Screening for RNAi targeting sequences against HBV. Using the siRNA target finder program of Ambion, nine potential sequences against HBV were selected. After two-step overlap PCR as shown in Fig. 1B, nine corresponding PCR products containing human U6 promoter, sense, loop, antisense and terminator sequence were obtained. The PCR products were co-transfected with the pHBV and pGL3-control plasmids into HepG2 cells and real-time PCR was used to quantify the HBV RNA and luciferase mRNA. The reduction of HBV RNA by these PCR products was calculated after normalizing the control luciferase mRNA level. Our results showed that these PCR products targeting HBV viral genes reduced the HBV RNA quantity from 17.2-95.6%, whereas the unrelated U6 PCR product (negative control NC) had no significant effect on the HBV RNA level (p=0.126). Among these PCR products, U6-shS2 and U6-shX1 were the most
effective and reduced HBV RNA by 95.6±3.51% and 92.3±1.52%, compared to that of the mock control (Fig. 1C). Thus, U6-shS2 and U6-shX1 were selected to construct siRNA-expressing HFV vectors for further study.

**Construction of siRNA expression HFV vectors and effective transduction.** Two selected siRNA expression cassettes were placed into the HFV vector to generate the HFVU6-siS2, HFVU6-siX1 and a dual siRNA expression vector HFVU6-siSX (Fig. 2A). The control plasmid pHFVU6-siNC was also constructed. All these HFV vectors contained the EGFP gene expressed from the HFV internal promoter (IP) which helped to monitor transduction efficiency. For determining the effective transduction titer of HFVU6 vectors, the 2.2.15 cells were transduced with the HFVU6-siNC vector at MOI of 5, 10, 30 and 50 for 7 days, and then the expression of EGFP was observed under an inverted fluorescence microscope. By counting the amount of cells with EGFP expression, it was shown that HFVU6-siNC transduced ~30% cells at MOI of 5, 70% at MOI of 10 and 95% at MOI of 30 and 50 (Fig. 2B). This same method was used to measure the titers of all the other HFVU6 vectors, and almost the same results were obtained (data not shown). Thus MOI of 30 was selected to transfect 2.2.15 cells for further study.

**Inhibition of HBV protein expression in 2.2.15 cells with transduced HFV vectors expressing siRNA.** 2.2.15 cells were infected with HFVU6-siS2, HFVU6-siX1, HFVU6-siSX and HFVU6-siNC at MOI of 30, respectively. The expression levels of HBsAg and HBeAg in culture media were determined with ELISA kits every 2 days post-transduction. As shown in Fig. 3A, three specific HFV siRNA vectors dramatically reduced the levels of HBsAg and HBeAg in the culture media, while the negative control vector had essentially no effect on HBV antigen expression. The reduction of HBsAg and HBeAg reached a peak on the 10th day post-transduction and remained level until day 14. HFVU6-siS2, HFVU6-siX1 and HFVU6-siSX reduced extracellular HBsAg and HBeAg levels by 90.5±1.4%, 83.6±0.7%, 92.4±2.7% and 84.8±1.7%, 79.5±2.1%, 88.7±1.9%, respectively, on the 10th day post-transduction.

**Immunofluorescence staining was employed to determine the expression levels of HBx protein in mock and transduced 2.2.15 cells on the 14th day post-transduction.** In the cells transduced with HFVU6-siX1 or HFVU6-siSX, HBx signals were barely detectable, while in the cells transduced with HFVU6-siS2 or HFVU6-siNC, the signals from HBx were quite strong and almost no different from the HBx signals in the non-transduced cells. This indicated that HFVU6-siX1 and HFVU6-siSX were able to efficiently inhibit the HBx protein expression while HFVU6-siS2 was ineffective (Fig. 3B, lane HBx), demonstrating the expected specificity.

As in all the HFV siRNA expression vectors containing the EGFP expression cassette, the EGFP signals were able to be used as a marker to label the transduced cells. In this study, 14 days post-transduction, the expression of EGFP was observed in >90% of the cells (Fig. 3B, lane EGFP). These data indicated that the HFV siRNA expression vectors infected the cells efficiently, and then induced stable and sequence-specific silencing of target genes of HBV.
Inhibition of HBV DNA in 2.2.15 cell culture media. To determine the effect of siRNAs produced by foamy virus vectors on viral DNA replication, the HBV DNA in 2.2.15 cell culture media was quantified by real-time PCR analysis. Quantitative assay revealed that HBV DNA levels decreased 4 days post-transduction. The greatest reduction was observed at 82.33±2.17%, 76.4±1.68% and 83.96±1.32% (p<0.05) on day 12 for the three HFV siRNA vectors, respectively. Meanwhile, HBV DNA levels were not significantly changed between untreated 2.2.15 cells and the cells transduced with control vector HFVU6-siNC (p>0.05) (Fig. 4).

Long-term anti-HBV efficacy of siRNAs produced by foamy virus vector. To assess the long-term anti-HBV efficacy of siRNAs produced by foamy virus vectors, the transduced 2.2.15 cells were continually cultured. Three months post-transduction (about 30 passages), total RNA was isolated and analyzed by semi-quantitative RT-PCR (Fig. 5A). Different sets of primers for cDNA synthesis were used to determine the levels of pgRNA, HBs and HBx mRNA in transduced cells. Results indicated that the levels of pgRNA mRNA were significantly decreased by the infection of HFVU6-siS2, HFVU6-siX1 and HFVU6-siSX, but not by HFVU6-siNC. HBs mRNA was decreased more strongly by HFVU6-siS2 and HFVU6-siSX, but slightly by HFVU6-siX1. Similarly, HBx mRNA was also decreased by HFVU6-siX1 and HFVU6-siSX. Notably, a reduced HBx mRNA level was observed in HFVU6-siS2 transduced cells. This was because pgRNA and HBs mRNAs also contained the HBx sequence at the 3' end. As the primers used to amplify HBx recognized not only HBx mRNA but also pgRNA and HBs mRNA, the total RNA templates for the HBx PCR reaction were reduced when HFVU6-siS2 reduced the levels of pgRNA and HBs mRNAs. This resulted in the decrease of HBx PCR products shown on the agarose gel. This explanation was supported by the previous immunofluorescence staining assay in which the HFVU6-siS2 had no effect on the HBx protein expression.

In addition, the inhibitory effects of the dual siRNA expression vector HFVU6-shSX on the levels of HBV pgRNA, HBs and HBx mRNA were greater than that of the single siRNA expression vector. This suggested that HFVU6-siS2 and HFVU6-siX1 may cooperate to inhibit HBV mRNA expression.

To further determine the effect of the stably expressed siRNAs on HBV DNA replication, intracellular HBV DNA was isolated and analyzed by Southern blotting. The levels of relaxed circular (RC), double-strand linear (DSL) and single-strand (SS) HBV DNA were clearly reduced in the cells transduced by HFVU6-siS2, HFVU6-siX1 and HFVU6-siSX but not in cells transduced by HFVU6-siNC. Density analysis further showed that HBV DNA replication intermediates were reduced by 76, 68 and 83%, respectively, compared to that of control (Fig. 5B).

Discussion

Recently, considerable attention has been directed toward the use of RNAi as a therapeutic method to treat a variety of diseases, especially viral diseases. Pioneer studies have demonstrated the possibilities of RNAi for treating serious viral diseases, including those caused by the human immunodeficiency virus, human corona virus, poliovirus, dengue virus, influenza virus and hepatitis C virus (17-23). In addition, a number of studies have shown that RNAi reduces...
the expression of HBV genes or/and attenuates the replication of HBV genome in cell culture or in transgenic mice (7,8,24-26). These studies have shed light on the development of new anti-HBV strategies.

To advance the application of RNAi in the inhibition of HBV, we employed a PCR-based RNAi strategy for a quick screening for effective anti-HBV siRNA sequences. A two-step overlap PCR strategy was utilized to generate a DNA template containing the U6 promoter and short hairpin DNA. The PCR products were directly co-transfected with a vector-bearing HBV genome into HepG2 cells, and HBV inhibition efficiencies were determined by real-time PCR. Compared to the traditional method, a multiple-step process which includes primer annealing, short hairpin sequence cloning into the RNAi expression vector, plasmid identification, amplification and transfection, this PCR-based strategy provides a rapid and easy assay for testing candidate siRNA sequences. By using this approach, we quickly identified two siRNA sequences (S2 and X1) having significant effects in the reduction of HBV RNAs.

Many studies have shown that viral vectors are good tools for effectively delivering foreign genes into cells. HFV is a member of the spumavirus subfamily of retroviruses (27). With its nonpathogenic nature, HFV has the advantage of being able to integrate into the host genome allowing for the stable expression of foreign genes, such as siRNAs. Therefore, in this study, HFV was employed to construct the siRNA expression vectors because it not only delivered the siRNA expression cassettes effectively but also allowed stable siRNA expression.

To study the long-term effects of selected siRNAs on HBV gene expression and viral replication, two single siRNA expression vectors HFVU6-siS2 and HFVU6-siX1, and a dual siRNA expression vector HFVU6-siSX were constructed and evaluated. These HFV siRNA vectors effectively inhibited multiple viral gene expression and viral DNA replication and most notably the reductions were stable for up to 3 months post-transduction. Several lines of evidence obtained from our study support the results. Firstly, ELISA and immunofluorescent staining experiments showed that viral proteins were significantly reduced in HFV siRNA vector transduced cells. Secondly, using RT-PCR, HBV RNAs were shown to be inhibited by these siRNA expression vectors. Finally, real-time PCR and Southern blotting showed that extracellular and intracellular HBV DNA were significantly reduced in HFV siRNA vector transduced cells.

Among the three vectors, HFVU6-siSX was a dual siRNA expression vector. It simultaneously expressed two siRNA expression cassettes effectively but also allowed stable siRNA expression. Thus, HFVU6-siSX was more efficient on the inhibition of gene expression and viral protein production compared to that of the single siRNA expression vectors. In addition, the use of the multiple-siRNA expression vector may overcome the obstacle of siRNA resistance. The fact that a single nucleotide mutation allowed the virus to escape the effect of a single specific siRNA has been reported in poliovirus and HIV infections (28,29). Recently, Wu et al also reported that a change of T to C at nucleotide 472 in an HBV clinical isolate resulted in siRNA resistance (26). Therefore, in this study we addressed such a potential problem by choosing targets in the relatively conserved viral sequences and by encoding two siRNAs that targeted different sites or genes simultaneously.

In conclusion, the PCR-based siRNA expression strategy provides a rapid and easy approach for testing candidate anti-HBV siRNA sequences and cloning selected siRNA expression cassettes into HFV vectors. RNAi based on the HFV vector achieved long-term inhibition of HBV gene expression and viral DNA replication in an HBV infection cell model. The strategy of constructing multi-siRNA expression vectors can enhance anti-HBV efficacy and overcome the evading mechanism of the virus.

References