Hepatitis B virus X protein activates human hepatic stellate cells through upregulating TGFβ1


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ABSTRACT. We investigated the effects of the hepatitis B virus X gene (HBV X) on the activation of human hepatic stellate cells (HSCs) and the possible mechanisms underlying the pathway. Recombinant plasmid pHBV-X-IRES2-EGFP was constructed and transfected into HL-7702 cells using a lipid-mediated method. Transfected cells were screened by G418, which detected stable expression of the X gene by reverse transcription (RT)-PCR and Western blot analysis, and named L02/x. Cells not subjected to G418-selection were analyzed to confirm the transient expression of the X gene and named L02/48x. Subsequently, L02/x and L02/48x, together with non-HBx-expressing cells, were co-cultured with HSCs in a non-contact transwell system. After 36 h of co-culture, the proliferation and migration of HSCs was detected using different cell counting methods. Finally, the mRNA and protein levels of α-SMA, Col I, and TGFβ1 in HSCs were detected by real-time PCR and western blot analysis. RT-PCR and Western blot analysis showed that L02/x and L02/48x cells can express HBV X gene mRNA and protein. Additionally, HSCs co-cultured with L02/x or L02/48x cells showed significantly higher proliferation and migration levels than
control groups. Real-time PCR and Western blot analysis showed that the mRNA and protein expressions of $\alpha$-SMA, Col I, and TGF$\beta$1 in HSCs co-cultured with HBx-expressing liver cells were higher than those in control groups. HBx protein activated HSCs in vitro, leading to increased proliferation and migration of HSCs and upregulation of $\alpha$-SMA and Col I. The TGF$\beta$1 gene may be involved in this pathway.

**Keywords:** Hepatitis B virus X (HBV X) gene; Gene activation; TGF$\beta$1; Hepatic stellate cells (HSC); Co-culture system

**INTRODUCTION**

Chronic hepatitis B virus (HBV) infection has been identified as a major risk factor in the development of cirrhosis and hepatocellular carcinoma (HCC), affecting more than 350 million people worldwide (Nebbia et al., 2012). The HBV X protein (HBx) is encoded by the HBV X gene, which is the smallest open reading frame of the HBV gene and is thought to play important roles in HBV-associated HCC by affecting gene transcription (Dewantoro et al., 2006; Matsuda and Ichida, 2009; Yip et al., 2011), DNA repair (Chen et al., 2008; Kim et al., 2010; Qadri et al., 2011), intracellular signal transduction (Chen et al., 2004; Ha and Yu, 2010; Hsieh et al., 2011), apoptosis (Gearhart and Bouchard, 2011; Knoll et al., 2011), and the cell cycle (Chen et al., 2008; Martín-Lluesma et al., 2008; Gearhart and Bouchard, 2011).

Recently, several studies have shown that HBx may be linked to HBV-induced fibrosis progression. Martín-Vílchez et al. (2008) showed that HBx protein induced paracrine activation of hepatic stellate cells (HSCs) in vitro and played a direct role in the development of liver fibrosis. Guo et al. (2009) found that HBx may facilitate liver fibrosis by promoting HSC proliferation. Additionally, these studies both revealed that the TGF$\beta$ gene may be involved in this pathway (Martín-Vílchez et al., 2008; Guo et al., 2009). However, the exact roles of the HBx protein in cirrhosis remain largely unknown and the molecular mechanisms of HBV-linked fibrosis remain unclear.

To explore the effects of the HBx protein on the development of HBV-associated cirrhosis, we constructed HL-7702 cells expressing the HBV X gene steadily or transiently. Additionally, HBx-carrying cells were co-cultured with HSCs in transwell systems to analyze the effect of HBx on HSC activation. Next, the expression of genes potentially related to these processes was detected to understand the mechanisms of HBx-associated fibrosis.

**MATERIAL AND METHODS**

**Construction of HBV X-expressing vector**

The HBV X DNA fragment was amplified by polymerase chain reaction (PCR) using the middle vector PUCmT-X, which was reserved by our laboratory as a template. The sequences of primers used for PCR were as follows: 5'-TAATCTCGAGATGGCTGCTAGGCTGTGCT-3' and 5'-GTCAGAATTCTTAATGGTGATGGTGATGGGGAGGAGTGAAAGTTGC-3'. PCR was carried out as follows: pre-denaturation at 94°C for 30 min and 35 amplification cycles (denaturation at 94°C for 30 s, annealing at 57°C for 30 s, and extension at 68°C for
Effects of HBx on HSCs

30 s). Next, PCR products were digested by XhoI and EcoRI and ligated into pIRES2-EGFP to construct the reconstituted plasmid pHBV-X-IRES2-EGFP. Its sequence was confirmed by restriction endonuclease digestion and direct DNA sequencing.

Cell culture and DNA transfection

HL-7702 cells were purchased from the Institute of Biochemistry and Cell Biology (Shanghai, China) and cultured in DMEM supplemented with 10% fetal bovine serum. Cells in the logarithmic growth were separately transfected with pHBV-X-IRES2-EGFP and pIRES2-EGFP plasmids using Lipo2000 transfection reagent, which was obtained from Invitrogen (Carlsbad, CA, USA). After 48 h, transfected cells were observed using a fluorescence microscope to evaluate transfection efficiency and the cells were divided into 2 groups. One group was selected using G418. After 2 weeks of selection, positive clones were isolated and further expanded, and these clones were referred to as L02/x and L02/ctr. Another group of cells was transfected transiently for 48 h without G418 selection and separately referred to as L02/48x and L02/48ctr. Corresponding control groups without transfection were referred to as L02/NC and L02/48NC, respectively.

RT-PCR

RNA from the 3 groups of cells (L02/x, L02/48x, and HL-7702) were extracted separately and reverse transcribed into cDNA according to manufacturer instructions of the MMLV reverse transcription kits, which were purchased from Promega (Madison, WI, USA). Using 2 mL RT products as the template, the X gene was amplified by PCR as follows: pre-denaturation for 2 min at 95°C and 40 amplification cycles (denaturation at 95°C for 20 s, annealing at 57°C for 15 s, and extension at 72°C for 20 s). Primers used to amplify the X gene and the control gene hACTB are shown in Table 1.

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence</th>
<th>Product (bp)</th>
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<td>HBV X-F1</td>
<td>CCGTGCTTGCCCTTCTCATC</td>
<td>105</td>
</tr>
<tr>
<td>HBV X-R1</td>
<td>ATCTCCTTCCCCAAACTCCTC</td>
<td></td>
</tr>
<tr>
<td>HBV X-F2</td>
<td>CTAAGGCTGTGCTGCCAACCTG</td>
<td>182</td>
</tr>
<tr>
<td>HBV X-R2</td>
<td>AGAAGGCACAGACGGGGAAGT</td>
<td></td>
</tr>
<tr>
<td>hACTB-F</td>
<td>TCCTTCTGGGGCATGGAAGT</td>
<td>208</td>
</tr>
<tr>
<td>hACTB-R</td>
<td>CAGGAGGAGCAATGATCTTGAT</td>
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</table>

Western blot

L02/x, L02/ctr, L02/NC, L02/48x, L02/48ctr, and L02/48NC cells were lysed in lysis buffer containing 50 mM Tris-HCL, pH 8.5, 150 mM NaCl, 0.2 g/L NaNO3, 0.1 g/L sodium dodecyl sulfate (SDS), 100 mg/mL phenylmethysulfonyl fluoride, 1 mg/mL aprotinin, 10 mL/L NP-40, and 5 g/L sodium deoxycholate. The products were centrifuged at 14,000 g for 15 min to remove cellular debris. Protein concentrations were determined by BCA methods according to the instructions of BCA protein assay kits. Equal amounts of each sample (50 μg) were subjected to 12% SDS-polyacrylamide gel electrophoresis (PAGE) and then transferred...
to a polyvinylidene difluoride (PVDF) membrane. The membrane was blocked in TBST (10 mM Tris-Cl, pH 7.4, 150 mM NaCl, and 10 mL/L Tween 20) containing 5% fat-free milk for 1 h and incubated overnight with the anti-HBx antibody (1:1000 dilution), which was obtained from Abcam (Cambridge, UK). The membrane was then washed 3 times for 15 min in TBST and incubated with anti-mouse immunoglobulin G for HBx. After washing, the bands were visualized by exposure to film with an ECL Western blot system from Zhongshan Biotechnology (Beijing, China).

**Construction of a co-culture system**

In a non-contact transwell co-culture system which was obtained from Corning Inc. (Corning, NY, USA), HSCs were cultured in the upper dishes, and 6 groups of live cells were separately cultured in the lower dishes in DMEM supplemented with 10% fetal bovine serum and 1% Glutamax. Next, HSCs co-cultured with L02/x, L02/ctr, L02/NC, L02/48x, L02/48ctr, or L02/48NC were referred to as HSC/x, HSC/ctr, HSC/NC, HSC/48x, HSC/48ctr, or HSC/48NC, respectively.

**Proliferation of HSCs**

After 36 h of co-culture, Cell Counting Kit (CCK)-8 reagents were added to the upper dishes about 10 μL per dish. Absorbance values at 450 nm were measured after 3 h. The absorbance value of the sample minus the value of blank was considered to be the final value. Each group was analyzed in triplicate.

**Migration of HSCs**

After 36 h of co-culture, moist swabs were used to erase the upper HSCs in the dishes. Next, the remaining cells were fixed with 4% paraformaldehyde for 15 min and dyed using crystal violet according to manufacturer instructions of crystal violet staining kits. The HSC migration was counted by measuring the absorbance at 570 nm using an absorbance microplate reader. Each sample was measured in triplicate.

**Real-time PCR**

The RNA from the 6 groups of co-cultured HSCs was extracted and reverse transcribed into cDNA. PCR primers for alpha-smooth muscle actin (α-SMA), type 1 collagen (Col I), and the control gene hACTB were synthesized according to their corresponding sequences (Table 2). The best annealing temperatures and amount of templates were confirmed by RT-PCR analysis. Real-time PCR was performed on a Mastercycler ep realplex PCR system (Eppendorf, Hamburg, Germany). The cDNA of each sample (1 μL) was amplified in a 20-μL reaction system containing 10 μL 2X SYBR Green Mix, and 1 μL primer mix. PCR conditions were as follows: predenaturation for 2 min at 95°C, followed by 40 cycles at 95°C for 15 s, 44-68°C for 30 s, and 68°C for 60 s. Data were collected and melting curves were analyzed. The relative expression level of each sample was calculated by comparing CT values with control groups. Each sample was run 3 times for each target gene.
Western blot

The processes were similar to those described above. Briefly, protein products were extracted from the 4 groups of HSCs, including HSC/x, HSC/ctr, HSC/NC, and HSCs without co-culture. Next, 50 μg samples were subjected to 10% SDS-PAGE and transferred to PVDF membranes. The membranes were blocked in 5% fat-free milk in TBST and incubated with the primary antibody against either α-SMA (1:1000 dilution), Col I (1:1000 dilution), or β-actin (1:4000 dilution) overnight at 4°C. α-SMA antibody was purchased from Abcam (Cambridge, UK). Col I alpha 1 (COL1A1) and β-actin antibodies were reserved by our laboratory. The membranes were then washed and incubated with the corresponding secondary antibodies. After washing, the bands were visualized by exposure to film with an ECL system. Each sample was run 3 times for each target gene.

RNA and protein expression of TGFβ1

Real-time PCR and Western blot analysis were used to detect the RNA and protein levels of TGFβ1. These procedures were similar to those described above. The primers required for real-time PCR are also described in Table 2. The primary antibody for TGFβ1 in Western blot was from our laboratory and diluted to 1:100.

Statistical analysis

Data are reported as means ± SD and statistical analysis was performed with independent samples t-test using the SPSS 17 software (SPSS, Chicago, IL, USA). P values <0.05 were considered to be statistically significant.

RESULTS

Construction of HBV X recombinant vector

A 465-bp HBV X gene fragment was amplified from PUCmT-X and sub-cloned into the expression vector pIRE2-EGFP to construct the recombinant vector pHBV-X-IRES2-EGFP. The sequence of the X gene in the plasmid was coincident with that reported previously as identified by restriction endonuclease digestion (Figure 1) and confirmed by DNA direct sequencing (Figure 2).

Table 2. Real-time PCR primers of α-SMA, Col I, TGFβ1, and control gene hACTB.

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence</th>
<th>Product (bp)</th>
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<tr>
<td>α-SMA-F</td>
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<td>161</td>
</tr>
<tr>
<td>α-SMA-R</td>
<td>TTGGTGATGATGCCATGTTCTAT</td>
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<tr>
<td>COL1A1-F2</td>
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<td>166</td>
</tr>
<tr>
<td>COL1A1-R2</td>
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<tr>
<td>TGFB1-F</td>
<td>ACAGAAATCTATGAACAGTTCAAGCA</td>
<td>197</td>
</tr>
<tr>
<td>TGFB1-R</td>
<td>GCTGAGTGATCCGCAAGGAA</td>
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</tr>
<tr>
<td>hACTB-F</td>
<td>TCCCTCTGAGGAGTATGCTTGAT</td>
<td>208</td>
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<tr>
<td>hACTB-R</td>
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Figure 1. Restriction endonucleases digestion analysis of the recombinant plasmid. The recombinant vector \( pHBV-X-IRES2-EGFP \) was digested by \( XhoI \) and \( BamHI \) and electrophoresed on a 1.5% agarose gel. A DNA fragment (X gene) of 470 bp was found in the digestion products of \( pHBV-X-IRES2-EGFP \), but not in the recombinant vector without digestion. It means that the recombinant plasmid of \( HBV \) X gene was constructed successfully. Lane 1 = 5000-bp DNA ladder; lane 2 = restriction endonuclease digestion products of \( pHBV-X-IRES2-EGFP \); lane 3 = \( pHBV-X-IRES2-EGFP \) without digestion.

Figure 2. Sequencing result of the recombinant plasmid \( pHBV-X-IRES2-EGFP \). The DNA sequence of \( pHBV-X-IRES2-EGFP \) was detected and compared with the \( HBV \) X gene through NCBI online. A similarity of 99% was found between them. The result indicated that the \( HBV \) X gene-expressing vector was constructed successfully.
Expression of the HBx gene

The pHBV-X-IRES2-EGFP and pIRES2-EGFP plasmids were transfected into HL-7702 cells. After 48 h, cells were divided into 2 groups. One group was selected by G418 for 2 weeks and named L02/x and L02/ctr, respectively. Another group of cells without G418-selection were named L02/48x and L02/48ctr, respectively. Non-transfected HL-7702 cells, as blank controls, were named L02/NC and L02/48NC. RT-PCR analysis showed that HBV X mRNA was detected in L02/x and L02/48x, but not in HL-7702 (Figure 3A). Western blot analysis showed protein expression of the HBV X gene in L02/x and L02/48x, but not in L02/ctr, L02/NC, L02/48ctr, or L02/48NC (Figure 3B). Thus, HBV X mRNA and protein were expressed steadily and transiently in L02/x and L02/48x, respectively.

Proliferation and migration of co-cultured HSCs

HSCs were then separately co-cultured with L02/x, L02/ctr, L02/NC, L02/48x, L02/48ctr, and L02/48NC. CCK8 kits were used to detect the proliferation of HSCs. The results showed that HSC/x or HSC/48x cells have significantly higher proliferation levels than the corresponding control groups (Figure 4A). Migration of HSCs was analyzed by crystal violet staining and OD_{570} detection. Statistical analysis showed that HSCs co-cultured with HBx-expressing live cells have significantly higher migration levels than the corresponding control groups (Figure 4B).

Expression of α-SMA, Col I, and TGFβ1

Real-time PCR revealed that the mRNA expression levels of α-SMA, Col I, and
**DISCUSSION**

HBV infection is associated closely with the development of liver fibrosis and HCC. Currently available evidence supports that HBx activity promotes the procession of HBV-linked HCC through a series of complex molecular mechanisms. However, the roles of HBx in the development of HBV-associated fibrosis remain unclear. Recently, Liu and Bai both reported that HBV affected the proliferation of HSCs *in vitro* (Liu et al., 2009; Bai et al., 2012). According to these studies, the PDGF-B/PDGFR-β signaling pathway and Col I may play important roles in this process (Liu et al., 2009; Bai et al., 2012). Additionally, Guo, Martín-Vílchez both found that HBx protein affected the activity of HSCs in the *in vitro* co-culture systems and revealed that TGFβ1 may be involved in this process (Martín-Vílchez et al., 2008; Guo et al., 2009). However, the exact mechanisms of HBx-leading fibrosis remain largely debated.
Effects of HBx on HSCs

To examine the relationship between HBx and liver fibrosis, HL-7702 cells carrying the HBV X gene were co-cultured with different groups of live cells. HSC/48NC and HSC/NC were used as references. Data are reported as means ± SD and analyzed with the t-test by the SPSS 17 software. The relative α-SMA mRNA expression levels of HSC/48x, HSC/48ctr, HSC/x, and HSC/ctr are shown as follows: 4.2488 ± 0.1180, 0.9774 ± 0.0280, 2.2715 ± 0.0592, 0.9076 ± 0.0109 (*P < 0.05 vs HSC/48ctr, *P < 0.05 vs HSC/48NC, *P < 0.05 vs HSC/ctr, *P < 0.05 vs HSC/x). The relative Col I mRNA expression levels of HSC/48x, HSC/48ctr, HSC/x, and HSC/ctr are shown as follows: 2.7898 ± 0.0509, 0.9505 ± 0.0076, 2.8484 ± 0.0525, 0.9615 ± 0.0038 (*P < 0.05 vs HSC/48ctr, *P < 0.05 vs HSC/48NC, *P < 0.05 vs HSC/ctr, *P < 0.05 vs HSC/x). The relative TGFβ1 mRNA expression levels of HSC/48x, HSC/48ctr, HSC/x, and HSC/ctr are shown as follows: 4.9145 ± 0.1365, 0.9186 ± 0.0369, 3.1461 ± 0.0705, 1.0046 ± 0.0040 (*P < 0.05 vs HSC/48ctr, *P < 0.05 vs HSC/48NC, *P < 0.05 vs HSC/ctr, *P < 0.05 vs HSC/x). The results revealed that α-SMA, Col I, and TGFβ1 mRNA expression levels in HSCs co-cultured with HBx-expressing live cells were significantly higher than those of the corresponding control groups. That was to say, HBx upregulated mRNA expression of α-SMA, Col I, and TGFβ1 in HSCs. Western blot analysis revealed that different protein expression levels of these four genes in HSC/x and control groups. Lane 1 = HSC/ctr; lane 2 = HSC/NC; lane 3 = HSC/x; lane 4 = HSCs without co-culture. α-SMA, Col I and TGFβ1 protein expression levels in four groups of HSCs. The protein expression levels of α-SMA in HSC/ctr, HSC/NC, HSC/x, and HSC are shown as follows: 0.6201 ± 0.0059, 0.6297 ± 0.0050, 1.7298 ± 0.0347, 0.6222 ± 0.0063. Col I protein expression levels in HSC/ctr, HSC/NC, HSC/x, HSC are shown as follows: 0.8385 ± 0.0014, 0.8407 ± 0.0011, 2.1185 ± 0.0064, 0.8392 ± 0.0015. TGFβ1 protein expression levels in HSC/ctr, HSC/NC, HSC/x, HSC are shown as follows: 0.8897 ± 0.0060, 0.8988 ± 0.0022, 2.1399 ± 0.0409, 0.9045 ± 0.0049. There were significant increases of α-SMA, Col I and TGFβ1 protein expression levels in HSC/x vs HSC/ctr, HSC/NC and HSC (P < 0.05). It means that HBx upregulated α-SMA, Col I and TGFβ1 protein expression levels in HSCs.

To examine the relationship between HBx and liver fibrosis, HL-7702 cells carrying the HBV X gene were co-cultured with HSCs, and then the proliferation and migration of HSCs were detected to determine the effects of HBx on the activity of HSCs. In this in vitro experiment, HL-7702, a normal human live cell line, was co-cultured with HSCs in non-contact transwell co-culture systems. This experimental design is similar to the in vivo environment, and may help to explain the impact of hepatitis B infection on HSCs. Our data revealed that expression of HBx indeed promoted the proliferation and migration of HSCs.

The activities of HSCs are tightly linked with the progression of liver fibrosis. When HSCs are activated, they show an increased capacity for proliferation, mobility, contractility, and synthesis of collagen and other components of the extracellular matrix (ECM) (Pinzani...
et al., 1998; Pinzani and Marra, 2001; Brandão et al., 2006). Excessive deposition of ECM is a characteristic of fibrogenesis, which effectively amplifies the fibrogenic response (Hui and Friedman, 2003; Parsons et al., 2007). Thus, increasing the proliferation and migration of HSCs indicate that HSCs are activated. In other words, expression of HBx activated HSCs and promoted the formation and development of liver fibrosis.

In addition, expression of α-SMA is also a significant sign of HSC activation (Brandão et al., 2006). Excessive synthesis of ECM components leads to increased expression of α-SMA as well as changes in the expression of L-type voltage-operated Ca\(^{2+}\) channels, which are known to mediate Ca\(^{2+}\) influx and regulate cellular contraction (Gasull et al., 2001; Ahmad and Ahmad, 2012). In our study, HSCs co-cultured with HBx-expressing liver cells showed increased mRNA and protein expression of α-SMA compared to those cells co-cultured with live cells without expressing HBx. Thus, HBx promoted the expression of α-SMA in HSCs and induced activation of HSCs, which is responsible for the development of hepatic fibrosis.

When HSCs are activated, excessive ECM products, particularly Col I, are produced and secreted (Pinzani and Marra, 2001; Ahmad and Ahmad, 2012). Thus, increased expression of Col I is an indicator of HSC activation. In this experiment, HBx indeed promoted the expression of Col I, which further confirmed that HSCs had been activated. Expression of the HBV X gene in HL-7702 activate the co-cultured HSCs, including the increased proliferation and migration of HSCs, the upregulation of α-SMA, resulting an increased plateau level of Col I and leading to the development of liver fibrosis.

The precise mechanisms responsible for the development of liver fibrosis remain unclear, but TGFβ is thought to participate in this pathway (Pinzani and Marra, 2001; Parsons et al., 2007). TGFβ1 is synthesized and secreted in a latent, biologically inactive form that can bind to the type II TGF-β receptor (TβRII) and lead to the phosphorylation and activation of the type I TGF-β receptor (TβRI). This activated TβRI kinase phosphorylates Smad2 or Smad3, which subsequently form a complex with Smad4 and migrate to the nucleus to regulate expression of target genes, including collagens (Bauer and Schuppan, 2001; Zong et al., 2012). In the process of hepatic fibrosis, TGFβ is the main mediator of fibrogenesis, which can promote activation of HSCs and induce sustained production of ECM components (Friedman, 2000; Pinzani and Marra, 2001; Bataller and Brenner, 2005; Moreira, 2007; Hayashi and Sakai, 2012). Our study confirmed that HBx protein upregulated the mRNA and protein expression of TGFβ1 in HSCs, leading to activation of HSCs and increased expression of Col I (Kisseleva and Brenner, 2007).

In summary, this study revealed that the expression of HBx activated HSCs by upregulating TGFβ1. Notably, HL-7702 cells and HSCs were co-cultured in non-contact transwell systems to simulate the in vivo environment. Thus, HBx protein plays roles via the paracrine pathway (Martin-Vilchez et al., 2008). Whether our conclusions apply to the in vivo environment requires further analysis.

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Specialist. We thank Nan-Hong Tang and Xiao-Qian Wang for providing technical support.

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