Abstract. Hepatitis B virus (HBV) has been revealed to be involved in the development of hepatocellular carcinoma. However, the mechanism remains to be fully elucidated. Smad-interacting protein 1 (SIP1) is a transcriptional repressor, which serves a pivotal role in cell metastasis. In the present study, the role of SIP1 in HBx-induced hepatocyte EMT and cancer aggressiveness was examined. It was found that HBV X protein (HBx) increased the expression of SIP1 and recruited it to the promoter of E-cadherin, resulting in depression of the transcription of E-cadherin. Histone deacetylase 1 was also found to be involved in the repressive complex formation. Furthermore, in an orthotopic tumor transplantation model in vivo, HBx promoted tumor growth and metastasis, whereas the knockdown of SIP1 attenuated the effect of HBx. These results indicate a novel mechanism for the development of HBV-related liver cancer.

Introduction

Liver cancer is a prevalent type of cancer in humans that ranks as the third leading cause of cancer-related mortality worldwide (1). Epidemiological studies have shown that chronic hepatitis B virus (HBV) infection is a major risk factor for liver cancer (2). As a member of the hepadnaviridae family, HBV is among the smallest of all known animal viruses. The integrated hepatitis B virion is a spherical particle, 42 nm in diameter, which is also known as the Dane particle (3). The HBV genome is a circular, partially double-stranded DNA of ~3,200 base pairs. Four overlapping open reading frames encode a viral envelope protein (pre-S1/pre-S2/S), a core protein (pre-C/C), a viral polymerase (P) and an HBV X protein (HBx) (4,5).

HBx, which is highly conserved within the species, is named after the encoding X gene as the amino acid sequence is not homologous to any known protein (6,7). It has been reported to be involved in the etiology of liver cancer through the transcriptional regulation of certain proto-oncogenes. However, the molecular mechanism underlying HBx-induced carcinogenesis remains to be fully elucidated. HBx affects hepatocyte proliferation and transformation by modulating signal transduction, such as the Wnt pathway. As a Wnt-regulated protein, E-cadherin is observed to decrease during epithelial-mesenchymal transition (EMT) and has been regarded as the most important factor in the progression of cancer to a more aggressive phenotype. Smad-interacting protein 1 (SIP1) is one of the most important transcriptional regulators for the expression of E-cadherin. In our previous study, it was found that SIP1 regulated HBV replication and expression (8); however, whether SIP1 is involved in HBV-related diseases has received limited attention.

In the present study, the role of SIP1 in HBx-induced hepatocyte EMT and cancer aggressiveness was examined. It was found that the exogenous expression of HBx resulted in hepatocyte EMT through interacting with SIP1. HBx combined with SIP1 and enhanced its ability to bind the promoter region of E-cadherin and repress its transcription. Histone deacetylase 1 (HDAC1) was also found to be involved in the repressive complex formation. Functional analysis demonstrated that the knockdown of SIP1 abrogated the effect of HBx on cell proliferation, migration and tumor aggressiveness. The present study is the first, to the best of our knowledge, to elaborate on the role of SIP1 in HBx-induced EMT and tumor aggressiveness.

Materials and methods

Plasmid DNA and lentivirus. The pcDNA3.1 and pcDNA3.1-HBx (pHBx) plasmids were prepared in the Key Laboratory of Molecular Biology of Infectious Diseases
Synthesis of small interfering (si)RNAs. The siRNAs targeting the HBx gene (siHBx) were designed against the conserved target region of the gene. Negative control siRNAs (siCont) were designed with scrambled sequences. All siRNAs were chemically synthesized by Sangon Corporation (Shanghai, China) and are shown in Table I.

Reagents and antibodies. The following primary antibodies were used in the present study: Rabbit anti-E-cadherin monoclonal antibody (Cell Signaling Technology, Inc., Beverly, MA, USA; cat. no. 3195), rabbit anti-N-cadherin monoclonal antibody (Cell Signaling Technology, Inc.; cat. no. 4061P), rabbit anti-SIP1 monoclonal antibody (Santa Cruz Biotechnology, Inc., Dallas, TX, USA; cat. no. sc-15393), mouse anti-β-actin monoclonal antibody (Santa Cruz Biotechnology, Inc.; cat. no. sc-20358), mouse anti-HDAC1 monoclonal antibody (GeneTex, Inc., Irvine, CA, USA; cat. no. GTX100513222) and mouse anti-β-actin monoclonal antibody (Boster Biological Technology, Ltd., Wuhan, China; cat. no. BM0627). The HDAC inhibitor trichostatin A (TSA) was obtained from Sigma-Aldrich; Merck KGaA (Darmstadt, Germany; cat. no. T1952). DAPI (Beyotime Institute of Biotechnology) and Lipofectamine 2000 (Invitrogen; Thermo Fisher Scientific, Inc.) were used according to the manufacturer's protocols.

Western blotting. Protein was extracted from the cells using a protein extraction kit (Kaiji, Jiangsu, China) according to the manufacturer's protocol. The protein concentration was measured with BCA protein assay reagent (Beyotime Institute of Biotechnology). Approximately 50 µg of protein was separated by 8% SDS-PAGE and blotted onto PVDF membranes (GE Healthcare Bio-Sciences, Pittsburgh, PA, USA). After being blocked with 5% non-fat milk for 1 h and blotted with the indicated primary antibodies (anti-E-cadherin, anti-N-cadherin, anti-vimentin, anti-SIP1 and anti-HDAC1 at 1:1,000 dilution; anti-Slug and anti-HBx at 1:500 dilution; anti-β-actin at 1:2,000 dilution) on a shaker at 4°C overnight, the membranes were incubated with secondary goat anti-mouse antibody (1:5,000 dilution Boster Biological Technology, Ltd., Wuhan, China; cat. no. BA1050) or goat anti-rabbit antibody (1:5,000 dilution; Boster Biological Technology, Ltd., Wuhan, China; cat. no. BM1054) for 1.5 h at 37°C. The levels of targeted protein in the cells were evaluated using the Bio-Rad electrophoresis documentation system (Gel Doc 1000, Bio-Rad Laboratories, Inc., Hercules, CA, USA).

Reverse transcription-quantitative PCR (RT-qPCR) analysis. Total RNA was isolated from the cells using TRIzol reagent (Invitrogen; Thermo Fisher Scientific, Inc.). The total RNA was then converted into single-stranded cDNA with a cDNA synthesis kit (Takara Bio, Inc., Japan). The reaction conditions were as follows: 37°C for 15 min and 85°C for 5 sec. Amplification of the targeted genes was performed on a CFX96TM Real-time PCR detection system (Bio-Rad Laboratories, Inc.) using SYBR Green PCR premix Ex Taq (Takara Bio, Inc.). The qPCR conditions were as follows: Initial denaturation at 95°C for 30 sec, followed by 40 cycles of 95°C for 5 sec and 60°C for 30 sec. The relative expression values of the targeted genes were calculated using the comparative Cq (2-ΔΔCq) method (9). The primers used for the qPCR were synthesized by Sangon Corporation (Shanghai, China) and are shown in Table I.

Immunofluorescent staining. Following rinsing with phosphate-buffered saline (PBS), the cells were fixed on coverslips with 4% paraformaldehyde, permeabilized with 0.5% Triton X-100, blocked with goat serum (Boster Biological Technology, Ltd., Wuhan, China; cat. no. AR0009) and subsequently incubated with the indicated primary antibody (anti-E-cadherin or anti-vimentin; 1:100 dilution) overnight at 4°C. Following washing three times, the cells were incubated with DyLight 549-Conjugated Goat anti-mouse secondary antibody (Abbkine Scientific Co., Ltd.; cat. no. A23310; 1:100 dilution) or DyLight 549-Conjugated Goat Anti-Rabbit Secondary Antibody (Abbkine Scientific Co., Ltd.; cat. no. A23320; 1:100 dilution) at 37°C for 1 h. The nuclei of the cells were stained

Cell culture and transfection. The HepG2 cells were maintained in Key Laboratory of Molecular Biology of Infectious Diseases (Chinese Ministry of Education, Chongqing Medical University). The plasmid expressing SIP1 (pcDNA4his/maxC-SIP1) was donated by Professor Janet E. Mertz (McArdrle Laboratory for Cancer Research, University of Wisconsin-Madison School of Medicine and Public Health). SIP1 short hairpin RNA (shSIP1) or non-targeting shRNA (shCont) were cloned into the pGMLV-SC5 plasmid vector provided by Genomeditech (Shanghai, China). The sequence of the SIP1-targeting shRNA was 5'-GAACAGACAGCTTACGA-3' and that of the shCont sequence was 5'-TGGTTCGACGTGTGCACGT-3'. The plasmids expressing shSIP or shCont were also packaged into a lentivirus. The pG3L3-Basic and pRL-TK plasmids were obtained from Promega Corporation (Madison, WI, USA, cat. no. E1751). The wild-type E-cadherin promoter (proE-cad-Luc) and the mutant E-cadherin promoter combined with a luciferase reporter (proE-cad-Luc-mEbox) were provided by Kumiko UiTei. The mutation sites of the reporters were provided by Kumiko UiTei. The mutation sites of the reporters were provided by Kumiko UiTei. The reporter sequences are shown in Table I.

Synthesis of small interfering (si)RNAs. The siRNAs targeting the HBx gene (siHBx) were designed against the conserved target region of the gene. Negative control siRNAs (siCont) were designed with scrambled sequences. All siRNAs were chemically synthesized by Genepharma (Shanghai, China) and are shown in Table I.
with 10 µg/ml DAPI at 37˚C for 10 min. The fluorescent images were then observed and analyzed by confocal fluorescence microscopy. Images were typically selected from those of three repeated experiments.

Transwell assays. The cells were suspended at a density of 10x10^5/ml in medium without FBS and then placed in the upper part of the Transwell unit with polycarbonate filters (Corning Costar, Cambridge, MA, USA). Medium containing 10% FBS was added to the lower wells of the chambers. Following overnight culture at 37˚C, the cells were fixed with 4% paraformaldehyde and stained with 0.1% crystal violet at 37˚C for 10 min. Three invasion chambers were used per treated group. The cells in the upper chamber were removed, washed with ddH_2O and dried in air. The cell migration abilities were quantified by counting the number of cells that had passed through the membrane and fixed on the underside.

Dual-luciferase assay. The cells were seeded at ~70% confluence in 24-well culture plates, co-transfected with a luciferase reporter vector promoter construct (pGL3-basic, proE-cad-Luc or proE-cad-Luc-mEbox) and pRL-TK (an internal control) with either pcDNA4hismaxC-SIP1, shRNA or a control vector using Lipofectamine 2000 (Invitrogen; Thermo Fisher Scientific, Inc.). The cells were then treated with TSA or DMSO. At 48 h post-transfection, the cells were collected and measured for luciferase activity on a GloMax microplate luminometer (Promega Corporation) using a dual-luciferase assay kit (Promega Corporation) according to the manufacturer's specifications. The firefly luciferase activity was normalized based on the Renilla luciferase activity. Assays were performed in triplicate and data are expressed as the mean ± standard deviation.

Treatments of the cells with TSA. The human hepatoma cell lines were plated at 30% confluence and cultured at 37˚C under 5% CO_2. TSA (500 µM) or DMSO (for control) were added and incubated for 48 h at 37˚C. During continued incubation, fresh culture medium was provided every 24 h. Extracts were collected from the cells following DMSO or TSA treatment and analyzed by western blotting.

Co-immunoprecipitation (Co-ip) assay. The sonicated cell lysates were added to 0.7 ml of RIPA lysis buffer (Beyotime Institute of Biotechnology), incubated with anti-HBx or anti-SIP1 antibodies on a shaker at 4˚C for 24 h, and incubated with protein A/G agarose (Beyotime Institute of Biotechnology) for 5 h. The immune-complexes were resolved by 10% SDS/PAGE, transferred onto PVDF membranes (GE Healthcare Life Science, Little Chalfont, UK) and probed with antibodies against HBx (1:500 dilution), SIP1 (1:1000 dilution) or HDAC1 (1000 dilution), as previously described in the western blotting protocol.

Chromatin immunoprecipitation (ChIP) analysis. The treated cells (at ~90% confluence) were cross-linked with 1% formaldehyde at 37˚C for 15 min and treated with glycine buffer (1:10; Beyotime Institute of Biotechnology) at 37°C for 5 min to quench the crosslinking reaction. Then, the samples were washed and resuspended in SDS lysis buffer (Beyotime Institute of Biotechnology) containing 1 mM PMSF. Following lysis on ice for 20 min, the samples were sonicated for 10 sec at 35 V with 30-sec spacing intervals for 10 min. Anti-SIP1 (1:200), anti-HBx (1:100) antibodies and immunoglobulin G (negative control), and protein A+G Agarose/Salmon Sperm DNA (Beyotime Institute of Biotechnology) were added for immunoprecipitation. The precipitated DNA was purified

| Table I. Primer and siRNA sequences. |
|--------------------------------------|
| Primer name                  | Sequence (5'–3')       |
| siRNA                        |                         |
| HBx siRNA sense            | AAGAGGACUCUCUGACUCUCAdTdT |
| HBx siRNA antisense        | UGAGAGUCCAAGACUGCCUCUDdT |
| Control siRNA sense        | UUCUCGCAACUGUGACGCUdTD |
| Control siRNA antisense    | ACUGUGACACGUUCGAGAAdTD |
| Chromatin immunoprecipitation |                         |
| Primer 1 forward            | AGGCAGGGATGCTACCTGAG    |
| Primer 1 reverse            | TGTTCTTGGCTCAGCCTACAC   |
| Primer 2 forward            | TAGAGGGTCACCCGCAGTCTAT  |
| Primer 2 reverse            | TCACAGGTTGCTTGGCATTTTC  |
| PCR                         |                          |
| E-cadherin forward         | TGGCCAGAAAATGAAAGAGG    |
| E-cadherin reverse          | GGTGATGTGGCAATGCTTCT    |
| Vimentin forward            | GCCAGGCAAGACGAGGAGT     |
| Vimentin reverse            | GGGTATCAACCCAGAAGGAGT   |
| SIP1 forward                | ATGGGGCAGAGCAGCAGAT     |
| SIP1 reverse                | GTGCAGTGCATGACCTCAGC    |
| β-actin forward             | CTCTTCCAGCCCTTCTCTCT    |
| β-actin reverse             | AGCACTGTGTGGCGTACAG     |
| Chromatin immunoprecipitation (ChIP) analysis. The treated cells (at ~90% confluence) were cross-linked with 1% formaldehyde at 37˚C for 15 min and treated with glycine buffer (1:10; Beyotime Institute of Biotechnology) at 37°C for 5 min to quench the crosslinking reaction. Then, the samples were washed and resuspended in SDS lysis buffer (Beyotime Institute of Biotechnology) containing 1 mM PMSF. Following lysis on ice for 20 min, the samples were sonicated for 10 sec at 35 V with 30-sec spacing intervals for 10 min. Anti-SIP1 (1:200), anti-HBx (1:100) antibodies and immunoglobulin G (negative control), and protein A+G Agarose/Salmon Sperm DNA (Beyotime Institute of Biotechnology) were added for immunoprecipitation. The precipitated DNA was purified
(OD 260/280=1.8-2.0) and subjected to PCR amplification. In total, Two pairs of primers specific to the human E-cadherin promoter were used for PCR analysis (Table I). The synthesized products were separated on a 1% agarose gel and visualized with the Gel Doc 1000 electrophoresis documentation system (Bio-Rad Laboratories, Inc.).

Cell proliferation assay. Cell Counting Kit-8 (CCK-8) assays were used to determine cell viability in the cell proliferation assays. In brief, the cells were cultured at a density of 5x10^4 in a 96-well plate. After transfection, the cells were incubated for 12, 24, 36 and 48 h, and 10 µl CCK-8 solution (Dojindo Molecular Technologies, Inc., Kumamoto, Japan) was added to each well and incubated for 2 h at 37°C. Absorbance at a wavelength of 490 nm was detected with a microplate reader. Each assay was performed three times in triplicate.

Cell apoptosis analysis. The cells were seeded in a 6-well plate at a density of 5x10^5 and transfected with the indicated plasmids for 48 h. The cells were then trypsinized, washed twice with PBS, resuspended in 95 µl of binding buffer and stained with 5 µl of Annexin V-FITC and 10 µl of propidium iodide (PI) working solution. Following incubating in the dark for 20 min, the cells were examined by flow cytometry.

Analysis of colony formation. For the clonogenicity analysis, at 24 h post-transfection, the transfected cells were seeded into 12-well plates at 500 cells (HepG2 cells) per well and maintained in complete medium for 2 weeks. The colonies were fixed with methanol and stained with crystal violet. The images were then observed using an optical microscope.

Tumorigenicity assays in nude mice. The animals were provided by the Laboratory Animal Centre of Chongqing Medical University (Chongqing, China). The use of animals complied with the institutional guidelines and was approved by the Ethics Committee of the Second Hospital Affiliated to Chongqing Medical University. Nude mice were used to establish a mouse xenograft model.18 mice (male, 4-6 weeks old) provided by the Laboratory Animal Centre of Chongqing Medical University (Chongqing, China). The use of animals complied with the institutional guidelines and was approved by the Ethics Committee of the Second Hospital Affiliated to Chongqing Medical University. The animals were housed under a humidity of 40-70% and a temperature of 20-22°C with a light:dark cycle of 12:12 h. The procedures were performed in accordance with the institutional guidelines and were approved by the Ethics Committee of the Second Hospital Affiliated to Chongqing Medical University. Nude mice were used to establish a mouse xenograft model.18 mice (male, 4-6 weeks old) were randomly divided into three groups, and housed under a humidity of 40-70% and a temperature of 20-22°C with a light:dark cycle of 12:12 h. The procedures were performed in accordance with the institutional guidelines and were approved by the Ethics Committee of the Second Hospital Affiliated to Chongqing Medical University.

In vivo metastatic model. The tumor masses were cut into 1x1 mm pieces and steeped in RPMI-1640 that had been supplemented with 100 U/ml of penicillin and streptomycin. Another 15 six-week-old nude mice were randomly divided into three groups. Subsequently, a piece of tumor from one tumor-bearing mouse was carefully implanted into the liver of one of the new mice to establish a hepatoma orthotopic transplantation metastatic model. To determine the time to detect the tumor metastases in vivo, a preliminary experiment was performed. After 8 weeks, the mice were sacrificed, and the livers, diaphragm and lungs were removed and prepared for subsequent histological examination. All animal experiments followed a blind and randomized animal study protocol.

Hematoxylin-eosin (HE) staining. Liver and diaphragm tissues were immersed in 4% paraformaldehyde for 4 h at room temperature, and then dehydrated through ascending alcohol series. Paraffin-embedded blocks were cut into 5-µm-thick sections, dewaxed in xylene at room temperature, and rehydrated through decreasing ethanol series. Then, Sections were stained with HE at room temperature for histological analysis.

Statistical analysis. The study results are representative of at least three independent experiments. The data are shown as the mean ± SD. Statistical analysis and graphical presentation were performed using the SPSS 16.0 (SPSS, Inc.) and GraphPad Prism software (version 6; GraphPad Software, Inc.). Student's t-test was used for all statistical analyses between groups. P<0.05 was considered to indicate a statistically significant difference.

Results

Influence of the ectopic expression of HBx on cell mobility and the expression of EMT-related proteins. EMT is an important precursor to the invasion and metastasis of a hepatoma tumor. To examine whether HBx induced EMT directly, the HepG2 cells were transfected with the pHBx and the expression levels of the EMT-related proteins were examined by western blot analysis and immunofluorescent staining methods. As shown in Fig. 1A, the expression of HBx resulted in downregulation of the epithelial marker E-cadherin and a concomitant increase in the expression of the mesenchymal markers vimentin and N-cadherin. The RT-qPCR assay confirmed that the transcription levels of E-cadherin decreased and those of vimentin increased (Fig. 1B). Similar changes in E-cadherin and vimentin were observed by immunofluorescent staining (Fig. 1C). To further confirm these results, the HepG2-X cells were also examined. Once a monoclonal strain was established, the HepG2-X cells lost their overlapping growth characteristics and exhibited a fibroblast morphology while the parental HepG2 cells maintained highly organized cell-cell adhesion (Fig. 1D). Western blot analysis confirmed the decreased expression of E-cadherin and increased expression of vimentin in the HepG2-X cells (Fig. 1E). These results were further confirmed by immunofluorescent staining (Fig. 1F). By contrast, when the overexpression of HBx in HepG2-X cells was knocked down by siRNA, the repressed levels of E-cadherin were restored and the expression of vimentin was reduced (Fig. 1G).

As increased cell migration is associated with EMT, Transwell assays were performed to detect cell mobility. The pHBx-transfected HepG2 cells demonstrated increased mobility in the Transwell assays compared with the pcDNA3.1 transfected cells. Similarly, increased cell mobility was observed in the HepG2-X cells compared with that in the parental cells (Fig. 1H).

Epigenetic repression of E-cadherin by HBx is associated with SIP1. As an essential EMT protein marker, the repression
of E-cadherin is normally required for cell migration and is associated with a poor cancer prognosis. The present study probed the molecular mechanisms through which HBx represses E-cadherin. In screening for the key regulators of the HBx-induced repression of E-cadherin in hepatoma cells, it was found that the transcription factor SIP1 was essential. Western blotting was used to investigate the expression of E-cadherin and several transcription factors implicated in EMT. Among these transcription factors, SIP1 exhibited increased expression in the pHBx-transfected HepG2 cells compared with that in the controls. SIP1 also exhibited a negative relationship with E-cadherin (Fig. 2A and B). A similar correlation was
found between the expression levels of E-cadherin and SIP1 in HepG2-X cells that stably expressed HBx (Fig. 2C). By contrast, the knockdown of HBx by siRNA transfection in the HepG2-X cells resulted in the reduced expression of SIP1 and concomitant upregulation of E-cadherin (Fig. 2D), indicating close links between SIP1, E-cadherin and HBx.

To further examine whether SIP1 is a crucial factor in HBx-induced EMT, the present study examined whether the knockdown of SIP1 reverted the EMT phenotypes. Following the suppression of SIP1 by shRNAs in HBx-expressing HepG2 cells, the E-cadherin levels were restored (Fig. 2E). Subsequent immunofluorescent staining was performed to confirm the results. As shown in Fig. 2G, the loss of SIP1 in HBx-expressing HepG2 cells resulted in the increased expression of E-cadherin and decreased expression of vimentin. The Transwell migration analysis showed that the promoting effects of HBx on cell migration were abrogated by the down-regulation of SIP1 (Fig. 2F). Together, these data suggested that
SIP1 may contribute to the epigenetic repression of E-cadherin mediated by HBx.

**SIP1 migrates to the E-cadherin promoter region to repress its expression with the cooperation of HBx.** As E-cadherin was downregulated at the transcriptional level following HBx introduction, it was hypothesized that HBx may directly regulate the promoter activity of E-cadherin. An E-cadherin promoter-luciferase reporter plasmid (proE-cad-Luc) was transfected into HepG2 cells to determine how HBx repressed the expression of E-cadherin. Consistent with the hypothesis, the overexpression of HBx significantly decreased the activity of the E-cadherin promoter in the HepG2 cells compared with that in the empty vector control or mock transfection cells (Fig. 3A).

As suggested in previous research, E-boxes within the human E-cadherin promoter are conserved among mammalian sequences (Fig. 3C). The two E-boxes (E-box1 and E-box3) that cover the proximal transcription start site have been shown to serve a crucial role in regulating E-cadherin by providing binding sites for SIP1 (10-12). To examine whether HBx regulates the expression of E-cadherin by enhancing SIP1 binding to E-cadherin promoter, ChIP assays were performed. The results revealed that HBx strengthened the binding of SIP1 to E-cadherin promoter fragments that spanned the E-boxes (Fig. 3B). To further elucidate the effect of this binding complex on the E-cadherin promoter, a luciferase reporter plasmid constructed with an E-cadherin promoter that had mutated E-boxes (mutated in both E-box1 and E-box3) was introduced (Fig. 3D). The results showed that the mutation of key nucleotides eliminated the HBx-induced inhibition of the reporter gene activity (Fig. 3E). By contrast, the knockdown of SIP1 in the pHBX-transfected HepG2 cells increased the wild-type E-cadherin promoter-driven luciferase activity (Fig. 3F). These data indicated that HBx may suppress the activity of the E-cadherin promoter by enhancing SIP1 binding to spaced E-boxes.

To elucidate whether there was a physical interaction between HBx and SIP1, Co-IP analysis of HBx-expressing HepG2 cells was performed. The results showed that SIP1 effectively bound to HBx in the HBx-expressing HepG2 cells (Fig. 3G). Subsequently, the subcellular distribution of HBx and SIP1 in HepG2-X cells were examined via immunofluorescence staining. As shown in Fig. 3H, SIP1 primarily localized in the perinuclear region of the cytoplasm; a lower level had localized in the same region in the HepG2-X cells (Fig. 4G). The subcellular localization of SIP1 and HDAC1 in immunofluorescence assays also suggested that SIP1 and HDAC1 localized in the same region in the HepG2-X cells (Fig. 4G).

**knockdown of SIP1 suppresses the oncogenic activity of HBx in HepG2 cells.** HBx is known to modulate cell growth, cell-cycle progression and migration. The present study subsequently examined whether SIP1 is involved in these processes. The CCK-8 assay indicated that the introduction of HBx significantly increased the cell growth rate compared with that in the control cells. When SIP1 was knocked down by shRNA, the increased proliferation rate of the HBx-expressing HepG2 cells almost returned to the original levels (Fig. 5A). These results suggested that SIP1 serves a vital role in HBx-regulated cell growth.

Dual staining ( Annexin V and PI) of the liver cancer cells was performed, followed by flow cytometric analysis to determine the effect of SIP1 and HBx on cell apoptosis. The Annexin V/PI assays indicated that there was a marked decrease in the apoptotic rate of the HBx-expressing HepG2 cells compared with that of the controls (Fig. 5C). However, the knockdown of SIP1 antagonized the effect of HBx and reverted the apoptotic rate.

Subsequently, colony formation assays were performed to further elucidate the effect of HBx and SIP1 on cell proliferation. As shown in Fig. 5B, HBx significantly enhanced HepG2 cell proliferation, whereas the knockdown of SIP1 by shRNA in HepG2-X cells resulted in a reduced proliferation rate. Taken together, these results showed that HBx is widely engaged in modulating the oncogenic activity of liver cancer cells in vitro and that SIP1 exerts an important influence in the process.
Figure 3. HBx recruits endogenous SIP1 to the E-cadherin promoter. (A) Dual-luciferase assay of human E-cadherin promoters in Mock treated, pcDNA3.1 or pHBX transfected HepG2 cells and HepG2-X cells. Data were normalized to the luciferase activity of cells treated with pcDNA3.1 and pGL3-Basic. Data are representative of at least three independent experiments. *P<0.05 HepG2 + pcDNA3.1 vs. HepG2 + pHBX; #P<0.05 HepG2 + pcDNA3.1 vs. HepG2-X. (B) ChIP primers were designed on E-cadherin gene regulatory regions. Distal primers correspond to downstream regulatory regions (1) of -1895 to -1707 nt of the E-cadherin gene. The ChIP primers (2) were designed adjacent to the TSS locations across the E-box region of the human E-cadherin promoter. pHBx-transfected HepG2 cell lysates were subjected to ChIP using an anti-SIP1 antibody. PCR was conducted using the indicated primer pairs. An empty vector and IgG served as external and internal negative controls. (C) Sequence homology of the consensus E-box in the E-cadherin promoter of mammalian. E-boxes 1, 3 and 4, CCAAT box and GC box are conserved regulatory elements, as shown in the diagram. The arrow indicates the putative TSS. (D) Mutations generated in the E-boxes carried the E-box 1 mutation CAGGTG → AAGGTA and E-box 3 mutation CACCTG → AACCTA. The wild-type E-cadherin promoter and promoter comprising two mutated E-boxes were cloned into a luciferase vector to construct the proE-cad-Luc and proE-cad-Luc-mEbox plasmids. (E) Dual-luciferase assay of E-cadherin promoter constructs with proE-cad-Luc or proE-cad-Luc-mEbox in pHBx- or pcDNA3.1-transfected HepG2 cells. **P<0.01. (F) Relative E-cadherin promoter activities in shSIP1/shCont and pcDNA3.1/pHBX treated HepG2 cells. *P<0.05; (shSIP1 + WT, vs. shCont + WT). (G) Co-immunoprecipitation in protein extracts of pcDNA3.1-transfected HepG2 cells and HBx-expressing HepG2 cells with anti-SIP1 or anti-HBx antibodies and western blot detection of HBx and SIP1, respectively. (H) Immunofluorescent staining of HepG2-X cells with anti-HBx and anti-SIP1 to show the subcellular co-localization of HBx and SIP1. DAPI was used to visualize nuclei. Scale bars=10 μm. SIP1, Smad interacting protein 1; HBx, hepatitis B virus X; pHBX, pcDNA3.1-HBx; sh, short hairpin RNA. TSS, transcription start site; WT, proE-cad-Luc; Mut, proE-cad-Luc-mEbox; ChIP, chromatin immunoprecipitation.
Figure 4. HBx recruits SIP1 and HDAC1 to the E-cadherin promoter to repress its expression. (A) Western blot analysis of E-cadherin and HDAC1 in pcDNA3.1 or pHBX transfected HepG2 cells +/- TSA. E-cadherin promoter activities in TSA-treated cells following transfection with (B) shSIP1 or (C) SIP1 expression plasmids. Results are reported as the relative luciferase activity, vs. activity of pGL3-Basic. *P<0.05, **P<0.01 (mean ± SD). (D) ChIP of lysates from HepG2 cells transfected with pHBX using anti-HDAC1 antibody. An empty vector pcDNA3.1 and protein G beads served as external and internal controls, respectively. (E) ChIP performed using HDAC1 antibody on the lysates of HepG2 cells treated with shSIP1/shCont and pcDNA3.1/pHBX. (F) Co-immunoprecipitation of HBx-expressing HepG2 cell-protein extracts with anti-SIP1 or anti-HBx antibodies and western blot detection of HDAC1, HBx and SIP1, respectively. (G) Immunofluorescent staining of HepG2-X cells with anti-HDAC1, anti-SIP1 and DAPI. The merged image showed HDAC1 and SIP1 co-localization in the nucleus. Scale bars=10 µm. SIP1, Smad-interacting protein 1; HBx, hepatitis B virus X; pHBx, pcDNA3.1-HBx; sh, short hairpin RNA; Cont, control; HDAC1, histone deacetylase 1; ChIP, chromatin immunoprecipitation; TSA, trichostatin A.
SIP1 is involved in HBx-induced tumorigenesis and liver cancer metastasis in vivo. To examine the role of SIP1 in HBx-induced tumorigenesis and liver cancer aggressiveness, the tumor formation of liver cancer cells expressing HBx was evaluated in tumorigenicity assays on athymic nude mice. The nude mice that were subcutaneously injected with the HepG2-X cells had a larger tumor volume and increased tumor weight compared with those received the parental HepG2 control cells (n=6 per group). When the expression of SIP1 was knocked down by shRNA interference in the HepG2-X cells, the tumor volume ratio and the tumor weight were reduced significantly (Fig. 6A-D). The effect of the knockdown of SIP1 was verified by western blotting (Fig. 6E).

To further determine the influence of SIP1 on the metastatic capacity of liver cancer cells caused by HBx in vivo, a hepatic orthotopic transplantation metastatic animal model was developed. The mice that were transplanted with the tumor masses from the HepG2-X cells carrying shCont developed more expansive liver tumors compared with those transplanted with tumors derived from the mock-transfected HepG2 cells (Fig. 6F). Visible intrahepatic metastatic tumors and the HE staining of liver tissue, with the exception of the orthotopically transplanted portion, indicated wider intrahepatic metastasis. The knockdown of SIP1 in HBx-expressing cells led to a significant decrease in liver tumor size and the scope of intrahepatic migration (Fig. 6F). Furthermore, tumor metastasis from the transplanted hepatic tumor to the diaphragm tissue was only detected in the HepG2-X tumor-transplanted mice. H&E staining of the diaphragm tissues verified that the metastatic tumor had the same morphology as the original hepatic tumors (Fig. 6H). Immunohistochemical analysis of the transplanted tumor confirmed the repressed levels of E-cadherin and increased levels of SIP1 following HBx introduction (Fig. 6G). Consistent with the in vitro results, the silencing of SIP1 efficiently restored the expression of E-cadherin in the tumors.

Discussion

Accumulated evidence supports HBV as an important cause of liver cancer by modulating different signal transduction pathways involved in EMT (13-17). Various transcriptional
regulators have been identified as important EMT mediators, including ZEB1, SIP1, Slug and Twist (18-20). As one of only two members of the vertebrate ZFH1 family, SIP1, also known as ZEB2, was initially found to bind to the MH2 domain of SMAD1. SIP1 was later revealed to interact with SMAD2, SMAD3 and SMAD5 (21,22) and to regulate Smad-dependent TGF-β signal transduction pathways. SIP1 was also reported to be implicated in embryonic development and cancer progression (23,24). However, whether SIP1 serves a role in HBV-induced liver cancer has not been discussed previously.
In the present study, it was found that ectopic HBx resulted in the increased expression of SIP1 and decreased expression of E-cadherin, leading to EMT change of the HepG2 cells. In addition, changes in the EMT protein markers induced by HBx were reversed by modulation of the expression of SIP1. As a member of the δEF1 family, SIP1 is characterized by a homeodomain with two clusters of highly conserved zinc fingers: An N-terminal cluster consisting of four zinc fingers (NFZ) and a C-terminal cluster containing three zinc fingers (CFZ) (23). SIP1 occupies promoter elements by NZF binding to one-spaced CACCT DNA sequences, including E-boxes (CACCTG), and CZF binding to the other. The binding of SIP1 to the two conserved E-boxes represses the activity of the E-cadherin promoter (25). The data obtained in the present study indicated that HBx suppressed the activity of the E-cadherin promoter by increasing the expression of SIP1 and enhancing its ability to efficiently bind to the E-cadherin promoter in HepG2 cells.

Previous studies have reported that HDAC1 is involved in the epigenetic modifications of tumor genes and tumor-suppressing genes in various types of cancer. The expression of HDAC1 in liver cancer is high (26). HDAC1 was selected as a candidate molecule for investigation in the present study. An increase of HDAC1 in HBx-expressing cells was observed, consistent with the former reports (27,28). Additionally, HDAC1 was found to form a triple complex with HBx and SIP1, which combined with the promoter of E-cadherin and repressed its transcriptional activity. ZEB1 has been reported to form a multi-protein complex with HDAC1 and HDAC2 to regulate the transcription of target genes (29,30). In the present study, SIP1 was also observed to partially co-localize with HBx and HDAC1 in subcellular sites. It appears that HBx recruits SIP1 and facilitates its binding to DNA, together with HDAC1. The present study focused only on HDAC1. However, other HDACs may be also involved in the regulation of ZEB2 (SIP1) and the investigation of other HDAC members is anticipated in the future.

The reduced expression of E-cadherin interrupts intercellular adhesion and conjunction, which facilitates the aggressiveness of tumors. HBx has been reported to promote liver cancer through influencing E-cadherin by several mechanisms, such as DNA methylation, Wnt, Snail and mSin3A (31-34). In the present study, SIP1 was identified as an important regulator in the HBx-induced modification of E-cadherin. The knockdown of SIP1 abrogated the promoting effect of HBx on cell proliferation, growth and migration. In subcutaneous transplantation mice and hepatoma orthotopic implantation mice, HBx promoted tumor growth, development and metastasis, whereas the sustainable knockdown of SIP1 efficiently attenuated the HBx-induced promotion of
tumorigenicity and aggressiveness. These data indicate a key role of SIP1 in the process of HBx-induced tumorigenicity and metastasis.

In conclusion, the present study demonstrated that SIP1 serves a pivotal role in HBx-induced liver cancer growth and metastasis. The study presents a novel mechanism for HBV-related liver cancer (Fig. 7) and may help to examine novel potential therapeutic approaches.

Acknowledgements

The authors thank Professor Janet E. Mertz (McArdle Laboratory for Cancer Research, University of Wisconsin-Madison School of Medicine and Public Health) for providing the pcDNA4hismaxC-SIP1 plasmids.

Funding

This study was supported by research grants from the National Natural Science Foundation of China (grant nos. 81672080 and 81873971) and a grant from Chongqing Health and Family Planning commission (grant no. 20141005).

Availability of data and materials

The datasets analyzed during the present study are available from the corresponding author on reasonable request.

Authors’ contributions

WXC initiated the project. DDL, PL, LD and BW designed and monitored the experiments. JJM, QH, JY, QFY and HC performed the experiments. YYY contributed to writing the manuscript. JC and WXC read and approved the final manuscript.

Ethics approval and consent to participate

The animals were provided by the Laboratory Animal Centre of Chongqing Medical University (Chongqing, China). The use of the animals complied with the institutional guidelines and was approved by the Ethics Committee of the Second Hospital Affiliated to Chongqing Medical University.

Competing interests

The authors declare that they have no competing interests.

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