In vitro Study on Hepatitis B Virus Infecting Human Choriocarcinoma JEG3 Cells and Its Mechanism

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Key Words
Hepatitis B · JEG3 · Placenta trophoblast · Infection · Southern blot · Real-time PCR

Abstract
Aim: To build a hepatitis B virus (HBV)-infected human trophoblast cell model in vitro and determine the mechanism of intrauterine HBV infection. Methods: Serum from hepatitis B-infected patients containing HBV DNA $>10^9$ was drawn, subsequently inoculated into human trophoblast cells in vitro (JEG3) and passage-cultured. The supernatants and intracellular HBV viral load of inoculated cells were tested by real-time PCR, and HBV DNA was determined by Southern blot. Results: From inoculation of the 1st passage JEG3 cells, the supernatant viral load of the 1st passage was seen increasing over time, which peaked at 120 h, whereas the HBV viral load was seen decreasing gradually in subsequent passages, and tested negative after the 6th passage. In addition, infected cells of HBV DNA were tested by Southern blot, and showed continual expression in the subsequent cell passages 1–5 while passage 6 was negative. HBsAg was tested as positive from different passages 1–5, and its concentration was also seen decreasing with each subsequent passage cultured until the 6th passage when it tested negative. Conclusion: HBV could infect human trophoblast cells (JEG3) in vitro, and it showed continual expression in subsequent cell passages 1–5.

Introduction

Hepatitis B is a global epidemic disease which poses significant dangers to human health [1]. China is a country with a high reported incidence of hepatitis B virus (HBV) infection, with about 120 million people carrying HBV. Over the years, prevention of HBV infection made a step forward when combined hepatitis B vaccines and hepatitis B immunoglobulin was introduced [2–5]; however, there are still vertical transmissions occurring at a reported rate of 5–15% each year [6]. Vertical transmission is one of the main routes of HBV transmission [7–9], while intrauterine transmission is a major factor which has resulted in the most HBV carriers, i.e. mother-to-child transmission.
Currently, the mechanism of intrauterine transmission has not yet been fully understood. Some researchers have studied the theory of placental infection and discovered that HBV could infect placental tissue [10, 11]. Hence, in vitro models of HBV infection have been our focus of study to date. Through utilization of a choriocarcinoma cell lines, JEG3, we have built an in vitro cellular model of HBV-infected placental trophoblast to aid us in observing the nature of viral infection in vitro in order to provide a laboratory basis for our study on HBV intrauterine transmission [12].

Materials and Methods

Reagents
Cell culture media, DME/F12 1:1 culture medium (Hyclon Inc.); fetal bovine serum (American Gibco Inc.); trypsin and antibiotics (penicillin and streptomycin, American Sigma Inc.); ELISA kit (Shanghai Kehua Bioengineering Co. Ltd.); anti-mouse antibiotics (penicillin and streptomycin, American Sigma Inc.); fetal bovine serum (American Gibco Inc.); trypsin and antibacterium reagent (Takara Biotechnology Dalian Co. Ltd.) were used.

JEG3 Cells and Cell Culture
JEG3 cell lines were obtained from Shanghai Institute of Cell Biology. The JEG3 cell culture DME/F12 1:1 culture solution (containing 12% fetal bovine serum, 100 U/ml penicillin, streptomycin, pH 7.4) was placed in an incubator under 5% CO₂ and containing 12% fetal bovine serum, 100 U/ml penicillin, streptomycin), and the culture solution, and sterile HBV(+) serum was added against 100 U/ml. When cell density reached higher than 90, 0.25% trypsin was used to digest passaged cells.

JEG3 Cells and HBV Infection
JEG3 cells were infected by HBV and were passage-cultured until cell density reached 70–80%, at which time the culture solution was discarded, replaced with serum-free DME/F12 1:1 culture solution, and sterile HBV(+) serum was added against 100 U/ml. The culture was then mixed gently, spread evenly onto cell surfaces and incubated at 37°C for 24 h. The infected fluid was then assimilated, washed with 0.01 mmol/l PBS 3 times for 3 min/wash. A negative control was designed by Takara Biotechnology (Dalian). The PCR product size was 148 bp. A standard curve was plotted based on HBV plasmid, the real-time PCR amplification curve and melt curve were determined at the end of the reaction, and the experimental results were analyzed by plotting a standard curve. Meanwhile, HBV DNA fluorescent quantitative PCR assay reagent (product of Shenzhen PG Biotechnology Co. Ltd.) was used for comparison testing [13].

HBV DNA Fluorescent Quantitative PCR
DNA was taken from cells and supernatants as templates, and HBV DNA was assayed respectively using our designed primer and quantitative reagents of PG Biotech’s HBV DNA fluorescent quantitative reagents. The total reaction volume was 25 μl. The reaction condition was set at 95° for 10 s, 95° for 5 s, 60° for 20 s for 40 cycles, and a melting curve was plotted at 65° for 15 s; and 94° for 2 min, 94° for 30 s, 55° for 45 s for 40 cycles. The CT value was recorded, HBV DNA plasmid was taken as a template, a standard curve was plotted and viral load was calculated from the standard curve obtained.

HBV DNA Southern Blot
For HBV DNA Southern blot detection, the HBV gene sequence GI:21326584 to design P52 probe primer was used: primer P1, 5’-AATCTCTTGTGACTTCTC-3’; and P2, 5’-CGTGTAAGTTGATG-3’ (nt 338–356). The PCR product size was 258 bp.

Supernatant HBsAg and HBcAg ELISA Assay of Infected Cells
HBsAg and HBcAg ELISA assay was performed on supernatants at each postinfection interval as well as in each passage, and a noninfected cell culture solution was used as a blank control. The sample OD value was 2.1 times higher than that of the negative control, which was interpreted as positive and vice versa.

Immunohistochemical Detection of HBsAg and HBcAg Expressions of Each Cell Passage
Fixated cell slides were washed with 0.01 m mol/l PBS 3 times for 3 min/wash and endogenous enzymes were inactivated with 3% H₂O₂. They were blocked with blocking buffer, primary antibody was added and they were kept at 4°C overnight. Products were washed with 0.01 m PBS 3 times for 3 min/wash, followed by the addition of egg albumin and bio-horseradish peroxidase conjugate, respectively, placed at 37°C for 15 min, and washed with 0.01 m mol/l PBS 3 times for 3 min/wash. A negative control was designed. Specimens were stained with DAB and subsequently with hematoxylin. Products were then treated with a gradient alcohol dehydration DMB coverslip using Permount. Obtained specimens were then observed under a light microscope and photographed.
**Results**

**HBV DNA Quantitative Assays of JEG3 Primary Cells**

Comparison testing results were obtained from our laboratory setups as well as PG reagents, which were basically identical. The DNA extracted from cells and the supernatants at variable time intervals after HBV infection of JEG3 primary cells were assayed by HBV DNA fluorescent quantitative PCR. The supernatant and intracellular HBV viral loads at each time interval showed an increasing trend over time (fig. 1).

**HBV DNA Quantitative Assays of JEG3 1st through 10th Passage Cells**

The infected cells were passage-cultured (1st through 10th passage, approx. every 96 h each subsequent passage), and the cultured cells and supernatants were collected at each passage, had their DNA extracted and were run through HBV DNA fluorescent quantitative PCR amplification assay. Cells were passage-cultured after being infected with HBV, and the HBV viral load of cells and supernatants at each passage was obtained. Both showed gradual decrement over the passages until the 6th passage when the HBV viral load was shown to be absent (fig. 2).

The HBV DNA quantitative PCR sample amplification curve can be seen in figure 3, the log-concentration can be seen in figure 4 and the amplification curve of the standard sample can be seen in figure 5. Real-time PCR product electrophoresis (148 bp) can be seen in figure 6.

**HBV DNA Southern Blot of JEG3 1st through 6th Passage Cells**

HBV DNA was obtained from JEG3 1st through 5th passage cells. Both showed gradual decrement over passages until the 6th passage when HBV viral load was shown to be absent (fig. 7).

**Supernatant HBsAg and HBeAg Detection in Infected Cells**

Supernatant HBsAg concentration in the early passages of infected cells was shown to increase over time (fig. 8), whereas HBeAg were all negative. Supernatant HBsAg could be detected in the primary to the 5th passages of cultured cells, and its concentration was shown to decrease over passages, becoming totally absent during the 6th passage (fig. 9). Supernatant HBeAg were shown to be negative in all cell passages.

**HBsAg and HBCAg Expression in Each Cell after Infection**

The SP immunohistochemistry technique was used to assay cell slides cultured at the 1st, 2nd, 3rd, 4th and 5th passages. During the 1st, 2nd, 3rd, 4th and 5th cell passages, in terms of HBsAg staining, brownish-yellow stains that were disseminated over the cytoplasm could be seen and assumed as positive. In terms of HBCAg staining, in the 1st, 2nd, 3rd, 4th and 5th infected cell pas-
Fig. 4. Log values of the HBV DNA standard sample initial copies.

Fig. 5. Amplification curve of the HBV DNA standard sample.

Fig. 6. JEG3 (1st through 7th passage after infection) supernatant (1–7) and cell (8–14) HBV DNA real-time PCR product electrophoresis (148 bp); (−) denotes normal cells, M denotes DNA maker.

Fig. 7. JEG3 (1st through 6th passage after infection) cell (1–6) HBV DNA Southern blot.

Fig. 8. Supernatant HBsAg concentration of 1st-passage-infected JEG3 cells at different time intervals.
release of HBsAg concentration and HBV load gradually increasing over time. Additionally, they were all positively correlated with each other, while intracellular HBV load was also shown to be increasing. This shows that HBV is able to infect trophoblast cells, and its viral load expression shows an increasing trend over a certain period of time. This fully demonstrated that, after 24 h of HBV infection, viruses are removed from cell surfaces through multiple rinses in general, but only the replication of HBV which has entered the cells may result in viral load and its secreted HBsAg increasing over time.

In 1986, Sureau et al. [22] reported that the use of closed-loop HBV DNA-infected hepatoma cell lines in vitro may promote the reproduction of intracellular HBV. However, the adoption of a transfection technique cannot mimic the natural infection status of HBV in terms of the regulatory effects of HBV towards cell membrane, infection process, cell-associated factors, etc. These have not yet been resolved. An HBV genome with a unique structure, the S gene in the S zone on the L chain (nt 155–835), transcribes and encodes HBsAg major protein under the control of the S gene promoter. The C gene promoter, CP(nt 1643–1849), is the key regulating factor of HBV’s replication and guides the transcription of 3.4- and 3.5-kb mRNA.

Some researchers have assumed that the structural integrity of HBV, particularly the high expression pre-SI
protein as well as regulatory factors required for viral replication, HBsAg concentration, etc., are the factors which promote the success of HBV infection into cultured cells in vitro [23]. HBV covalently closed-loop DNA (cccDNA) is the synthetic template of intermediate mRNA and pre-genome RNA in HBV genome replication, and is a key factor which leads to persistent HBV infection [24–27]. In order to justify the presence of HBV replication in placental trophoblast cells, it is necessary to perform HBV cccDNA detection during the construction of the in vitro HBV-infected trophoblast cell model on which we are currently concluding this corresponding study. With the joint efforts of all researchers, we believe that these studies will present useful evidence for science and provide us with a better understanding of the mechanisms of intrauterine HBV infection.

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