Rapid quantification of hepatitis B virus DNA by real-time PCR using efficient TaqMan probe and extraction of virus DNA

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Abstract

AIM: To rapidly quantify hepatitis B virus (HBV) DNA by real-time PCR using efficient TaqMan probe and extraction methods of virus DNA.

METHODS: Three standards were prepared by cloning PCR products which targeted S, C and X region of HBV genome into pGEM-T vector respectively. A pair of primers and matched TaqMan probe were selected by comparing the copy number and the Ct values of HBV serum samples derived from the three different standard curves using certain serum DNA. Then the efficiency of six HBV DNA extraction methods including guanidinium isothiocyanate, proteinase K, NaI, NaOH lysis, alkaline lysis and simple boiling was analyzed in sample A, B and C. Three standards were prepared by cloning PCR products which targeted S, C and X region of HBV genome into pGEM-T vector respectively. A pair of primers and matched TaqMan probe were selected by comparing the copy number and the Ct values of HBV serum samples derived from the three different standard curves using certain serum DNA. Then the efficiency of six HBV DNA extraction methods including guanidinium isothiocyanate, proteinase K, NaI, NaOH lysis, alkaline lysis and simple boiling was analyzed in sample A, B and C by real-time PCR. Meanwhile, 8 clinical HBV serum samples were quantified.

RESULTS: The copy number of the same HBV serum sample originated from the standard curve of S, C and X regions was $5.7 \times 10^4$/mL, $6.3 \times 10^4$/mL and $1.6 \times 10^5$/mL respectively. The relative Ct value was 26.6, 31.8 and 29.5 respectively. Therefore, primers and matched probe from S region in combination with NaOH lysis is a simple, rapid and accurate method for quantification of HBV serum DNA.

CONCLUSION: Real-time PCR based on optimized primers and TaqMan probe from S region in combination with NaOH lysis is a simple, rapid and accurate method for quantification of HBV serum DNA.

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Key words: Hepatitis B virus; Serum DNA; Real-time PCR; Extraction method

INTRODUCTION

Hepatitis B virus (HBV) is a human hepadnavirus that causes acute and chronic hepatitis and hepatocellular carcinoma[1]. Although an effective vaccine has been available for two decades, an estimated 350 million people worldwide are chronically infected with HBV[2]. The conventional ways available for curing this disease are not very efficient. A significant proportion of chronic infections terminate in hepatocellular carcinoma, leading to more than one million deaths annually[3]. The diagnosis and clinical monitoring of HBV infection are based on the detection of viral antigens, antibodies to viral proteins, and circulating viral genome (HBV DNA)[4-6]. There is increasing evidence that measuring the level of HBV DNA in serum is useful in monitoring the efficacy of antiviral therapy, detecting the occurrence of drug-resistant mutants and relapse after discontinuing antiviral therapy[7-9]. An accurate quantitative assay for serum HBV
DNA may monitor residual viral load during treatment and allow the timely detection of flares of viral replication that accompany the emergence of variants.

In China, enzyme-linked immunosorbent assay (ELISA) is still a main detection method for HBV infection, but ELISA results neither efficiently reflect serum viral load or hepatitis activity nor monitor the efficacy of antiviral treatments. In recent years, real-time PCR detection assays based on SYBR Green I\[10,11\] and TaqMan probe\[12-19\] have been proposed for quantification of HBV DNA in serum. In the former assay, SYBR Green I can specifically bind to double-stranded DNA (dsDNA) rather than to single-stranded DNA (ssDNA). Fluorescence is monitored once each cycle after product extension and increases above background fluorescence at a cycle number that is dependent on the initial template concentration. Unfortunately, the dye detects all dsDNAs, including primer dimer and other nonspecific products.

The real-time PCR method based on the TaqMan probe uses a dual-labeled fluorescent probe containing a reporter dye that is quenched by a second fluorescent dye\[20\]. Hydrolysis of the probe by polymerase exonuclease activity separates the reporter from the quencher as the amplification proceeds, the fluorescence signal from the reporter increases cumulatively. The cycle at which fluorescence exceeds background, known as the threshold cycle, is inversely related to the initial copy number, thus allowing quantitative analysis.

The efficacy and accuracy of real-time PCR largely depend on the primers and probe\[21-23\], and are also related to the extraction method of HBV DNA\[24,25\]. A simple, rapid, efficient method for DNA extraction is crucial to the success of real-time PCR and the subsequent analysis. We describe here a rapid, convenient real-time quantitative assay for serum HBV DNA which combines optimized primers and probe with simple HBV DNA extraction method. The assay was designed to enable accurate quantification of clinical serum samples, which makes it a useful clinical test to monitor serially the efficacy of antiviral therapy.

**Materials and Methods**

**Preparation of standards**

Serum HBV DNA was extracted by NaOH lysis as previously described\[26\]. In brief, 50 µL of HBV positive serum was mixed with an equal volume of 0.4 mol/L NaOH and then the mixture was incubated at 80°C for 10 min, followed by centrifugation for 30 s at 15000 × g. After that the supernatant was carefully transferred to a new microcentrifuge tube and supplemented with 25 µL of 0.4 mol/L Tris-HCl (pH7.5). Two microliters of HBV DNA was used as a template in PCR.

Three pairs of primers were designed by software Beacon Designer 2.1 in the conserved region of S, C and X gene respectively and synthesized by Sangon Co., Ltd, Shanghai. The sequences of each pair of primers were as follows: RSU (5’-AGAATTCCTCACAATACCGCAGA GT-3’) and RSL (5’-CACACGGGTAGTCCCCCTAGA A-3’), RCU (5’-GTCTTTCGAGTGAGTGGATTG-3’) and RCL (5’-CGGCGATGTGACCTTGTTG-3’), RXU (5’-ACTCCGCTTTGCTGCCTTCT-3’) and RXL (5’-CATTCCGTCGGCGTTCAC-3’).

Three amplified products were cloned into the pGEM-T vector (Promega, USA), and then the recombinant HBV plasmids pGEM-S, pGEM-C and pGEM-X were serially diluted from 10⁷ to 10³ copies/µL after identification by PCR. One microliter of each diluted recombinant plasmid was used as a standard PCR template.

**TaqMan probes**

Taqman probes targeting S, C and X genes were designed by software Beacon Designer 2.1 and synthesized in Sangon Co., Ltd, Shanghai. The sequences of PS, PC, and PX were 5’FAM-AGACTCGTGGTGGACCTCTCAAT-TAMARA3’, 5’FAM-TCCCTAGAAGAAGAACTCCCTGCGCTC- TAMARA3’, and 5’FAM-CCGGACCGTGTTGCA CTTCGCTT-TAMARA3’, respectively.

**DNA extraction**

Six methods (including guanidinium isothiocyanate, proteinase K, NaI, NaOH lysis, alkaline lysis, as well as boiling) were used to extract serum HBV DNA from three serum samples\[24-28\]. Two microliters of HBV DNA was used as a template in real-time PCR.

In the methods of GuSCN, proteinase K and NaI lysis, protocol after the lysis process was the same, though components of lysis were different. GuSCN lystate consisted of 1 mol/L Tris-HCl (pH8.0), 0.5 mol/L EDTA (pH8.0), 100 g/L SDS, 200 mg/L proteinase K\[24\]. Proteinase K lystate contained 10 mmol/L Tris-HCl (pH8.0), 10 mmol/L EDTA (pH8.0), 0.5% SDS, 150 mmol/L NaCl and 200 mg/L proteinase K\[27\]. NaI lystate buffer included 6 mol/L NaI, 0.5% SDS, 26 mol/L Tris-HCl (pH8.0) and 13 mmol/L EDTA (pH8.0)\[28\]. First, 100 µL lystate was added into an equal volume serum, then the mixture was incubated at 37°C for 1 h, 55°C for 2 h and 60°C for 15 min in GuSCN, proteinase K and NaI lysis methods respectively. After that, 200 µL chloroform/isooamyl alcohol (24:1) was added to extract products and the supernatants were removed into new eppendorf tubes after centrifugation at 15000 × g for 15 min, followed by alcohol precipitation and further washing\[26,25,27\]. At last, naturally dried DNA precipitates were dissolved in 30 µL 0.1 × TE.

The easily manipulated alkaline lysis and boiling extraction were similar to the above NaOH lysis, except for the lystate. The lystate used in alkaline lysis was composed of 1 mol/L NaOH, 2 mol/L NaCl and 0.5% SDS\[26\], while the boiling extraction consisted of an equal volume of PBS\[28\].

**Optimization of primers and probes**

Real-time PCR amplification was performed in 20 µL...
reaction mixture containing 1 μL standard DNA or 2 μL isolated serum HBV DNA, 2 × PCR reaction buffer [100 mmol/L Tris-HCl (pH8.3), 100 mmol/L KCl, 7.0 mmol/L MgCl\textsubscript{2}, 400 μmol/L each of the deoxynucleotide triphosphates (dNTP), 1U hot star DNA polymerase Takara, Japan], 5pmol of each pair of primers and 2.5 pmol of corresponding TaqMan probe. All amplification reactions were performed in triplicate. After preparation of the reaction mixtures in 96-well plates, amplification was performed as follows: initial denaturation at 94°C for 3 min, followed by 40 cycles at 94°C for 20 s, 60°C for 40 s. Fluorescence readings were recorded at 60°C in each cycle. Results were analyzed by the software iCycler\textsuperscript{TM} iQ 3.0a provided with the iCycler system (Bio-Rad, USA).

**Evaluation of six DNA extraction methods**

To evaluate HBV DNA extraction methods, RSU, RSL primers and PS probe were used in real-time PCR. Amplification and analysis were performed as above.

**Quantification of HBV viral DNA in sera by real-time PCR**

NaOH lysis was performed to extract serum HBV DNA from 8 clinical serum samples. The selected RSU, RSL and PS were used in real-time PCR. The procedures of real-time PCR and standard curve analysis were described as above.

**RESULTS**

**Linear standard curve and optimization of primers and TaqMan probes**

Three recombinant plasmids pGEM-S, pGEM-C and pGEM-X were used as standards to set up the real-time protocol and construct the standard curve. Forty cycles of amplification allowed us to obtain linear quantification between 1 × 10\textsuperscript{3} and 1 × 10\textsuperscript{7} copies/μL per reaction (Figure 1). The highest copy number (5.7 × 10\textsuperscript{4}/mL) and the lowest Ct value (26.6) of the same HBV serum samples were observed from the standard curve originated from S region (Table 1). The lowest copy number (6.3 × 10\textsuperscript{2}/mL) and the highest Ct value (31.8) were displayed from the

<table>
<thead>
<tr>
<th>Standard curve</th>
<th>Ct</th>
<th>Copy number (/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>S</td>
<td>26.6 ± 0.061</td>
<td>0.2</td>
</tr>
<tr>
<td>C</td>
<td>31.8 ± 0.073</td>
<td>0.2</td>
</tr>
<tr>
<td>X</td>
<td>29.5 ± 0.357</td>
<td>1.2</td>
</tr>
</tbody>
</table>

CV: Coefficient variation.
standard curve of C region. The highest copy number from S region was about 90.5 and 35.6 multiple to the C and X regions. Hence, RSU, RSL primers and PS probe were chosen for further analysis because of the efficient amplification.

Different DNA extraction methods

To explore a simple and fast method for serum HBV DNA extraction, six different methods were compared by real-time PCR in combination with selected primers and TaqMan probe from S region. The copy number and Ct values of three HBV serum samples on the standard curve (Figure 2 and Table 2) showed that NaOH lysis and boiling were predominately advantageous over the other methods. Adversely, alkaline lysis failed to detect any HBV DNA in all samples. There was no significant difference among the other three methods.

Quantitation of HBV viral DNA in sera by real-time PCR

Table 3 indicated the copy number and Ct values of 8 positive HBV serum samples based on the optimized primers and TaqMan probe as well as DNA extraction. Their SD and CV unfolded that quantification of hepatitis B virus DNA in triplicate by real-time PCR was reliable and accurate.

**DISCUSSION**

Detection and analysis of PCR products should be carried out simultaneously with temperature cycling during amplification. If the fundamental properties of DNA, such as product size, quantity, sequence, or melting profile can specifically identify the products during PCR, no further analysis is required.

Real-time PCR is a powerful diagnostic tool capable of rapidly generating reliable and reproducible results with reduced risks of cross contamination[29]. The amplification efficiency can be detected with fluorescent probes. All steps are accomplished automatically by computer except for sample preparation, which can monitor PCR reaction timely.

To make our assay more sensitive and efficient, we...

<table>
<thead>
<tr>
<th>Method</th>
<th>Copy number/mL and Ct value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample A</td>
<td>Sample B</td>
</tr>
<tr>
<td>Guanidinium isothiocyanate</td>
<td>$1.41 \times 10^5 \pm 1.23 \times 10^4$</td>
</tr>
<tr>
<td>Proteinase K</td>
<td>$2.99 \times 10^5 \pm 2.88 \times 10^4$</td>
</tr>
<tr>
<td>Nal</td>
<td>$3.05 \pm 0.19$</td>
</tr>
<tr>
<td>NaOH lysis</td>
<td>$2.51 \times 10^5 \pm 2.91 \times 10^4$</td>
</tr>
<tr>
<td>Boiling</td>
<td>$3.49 \times 10^8 \pm 2.72 \times 10^7$</td>
</tr>
<tr>
<td>Alkaline lysis</td>
<td>N/ N</td>
</tr>
<tr>
<td>Boiling</td>
<td>$2.14 \times 10^9 \pm 2.16 \times 10^8$</td>
</tr>
<tr>
<td></td>
<td>$23.9 \pm 0.10$</td>
</tr>
</tbody>
</table>

N: Negative.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Ct</th>
<th>Copy number/mL</th>
<th>Mean ± SD</th>
<th>CV (%)</th>
<th>Mean ± SD</th>
<th>CV (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>25.4 ± 0.24</td>
<td>0.9</td>
<td>1.4 \times 10^7 ± 2.38 \times 10^6</td>
<td>17.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>31.9 ± 0.20</td>
<td>0.6</td>
<td>1.4 \times 10^7 ± 1.87 \times 10^7</td>
<td>13.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>31.5 ± 0.52</td>
<td>1.7</td>
<td>1.8 \times 10^7 ± 2.32 \times 10^7</td>
<td>12.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>37.2 ± 0.14</td>
<td>0.5</td>
<td>3.5 \times 10^7 ± 3.73 \times 10^7</td>
<td>10.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>34.0 ± 0.27</td>
<td>8.1</td>
<td>6.4 \times 10^6 ± 6.68 \times 10^6</td>
<td>9.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>32.2 ± 0.52</td>
<td>0.2</td>
<td>1.1 \times 10^7 ± 4.45 \times 10^6</td>
<td>4.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>27.0 ± 0.23</td>
<td>0.8</td>
<td>4.6 \times 10^7 ± 7.22 \times 10^7</td>
<td>15.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>33.9 ± 0.12</td>
<td>0.4</td>
<td>3.1 \times 10^7 ± 2.76 \times 10^7</td>
<td>8.9</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

CV: Coefficient variation.
optimized three pairs of primers and probes targeting the conserved S, C and X regions of HBV genome respectively. The highest copy number and Ct value of HBV serum samples were observed on the standard curve originated from S region. This indicated that the sensitivity of real-time PCR can be improved by adjusting primers and probe.[23]

HBV DNA extraction from serum is a key step in real-time PCR, as it directly affects the accuracy of quantification of viral load. NaOH lysis showed that the highest copy number of HBV serum samples A, B, and C was $3.49 \times 10^7$ copies/mL, $2.08 \times 10^7$ copies/mL and $4.40 \times 10^6$ copies/mL, respectively. Boiling was as effective as NaOH lysis in DNA extraction from HBV serum samples A, B, and C. The copy number of HBV serum samples A, B, and C detected by boiling was $2.14 \times 10^7$ copies/mL, $1.83 \times 10^7$ copies/mL, and $1.48 \times 10^6$ copies/mL, respectively. However, boiling was unsuitable for DNA extraction from serum with high viscosity.[24]

Compared with the methods of NaOH lysis and boiling, guanidine isothiocyanate, proteinase K and NaI decreased the copy number of HBV serum samples A, B and C to approximately $10^5$ to $10^3$ copies/mL. Proteinase K lysis was a bit better than the other two methods. In all three methods, the lysis was mixed with serum and extracted with chloroform/isoamyl alcohol (24:1), followed by precipitation and washing with absolute alcohol. A large quantity of DNA was lost due to the complicated process of these methods. DNA extraction with alkaline failed to detect serum DNA in our experiment, possibly due to the high concentration and pH value. On the contrary, $0.4 \text{ mol/L NaOH}$ lysis followed by neutralization with $0.4 \text{ mol/L Tris-HCl (pH7.6)}$ showed excellent efficacy because of a moderate alkaline concentration and lower pH value which may significantly improve the efficiency of PCR amplification.

In conclusion, real-time PCR based on the optimized primers, probe and DNA extraction is a simple, accurate, specific and sensitive method for the measurement of HBV viral load in serum. NaOH lysis for HBV serum DNA extraction is rapid, simple and efficient, making the assay suitable for handling a large number of clinical HBV serum samples. Since there are about 200 million HBV carriers in China, doctors need to know the HBV status of patients before starting any medical treatment. This assay may be especially useful for monitoring the therapeutic effects in chronically infected patients on antiviral therapy.

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