Rapid quantification of hepatitis B virus DNA by direct real-time PCR from serum without DNA extraction

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The purpose of this study was to quantify hepatitis B virus DNA by direct real-time PCR from serum without the need for DNA extraction. Crossing point (Cp) values were determined automatically using the second derivative maximum mode. Since serum samples from patients are inevitably haemolysed, lipaemic or icteric, the interference of endogenous substances from the serum in real-time PCR was evaluated. The result showed that, although serum protein quenched the intensity of fluorescence, the Cp value adopted to calculate the quantity of DNA copies remained unchanged. Importantly, real-time PCR from serum with or without DNA extraction reached a high level of concordance. This direct serum PCR method without the DNA extraction and gel electrophoresis allows for substantial labour and cost savings. In addition, it is also suitable for rapid DNA quantification during clinical diagnosis.

INTRODUCTION

Although classical PCR-based assays have increased the sensitivity of analyses, their application in clinical practice is hindered by technical complexity, susceptibility to contamination and variable reproducibility (Kaneko et al., 1990; Quint et al., 1995). Recent advances in PCR technology allow continuous measurement of the fluorescence emitted during amplicon production after each PCR cycle. This is achieved using a variety of different fluorescent chemistries that correlate PCR product concentration to fluorescence intensity (Higuchi et al., 1993). Reactions are characterized by the time point (or PCR cycle) where target amplification is first detected. This value is usually referred to as the cycle threshold (Ct) or crossing point (Cp), the time at which fluorescence intensity is significantly greater (usually ten times the SD of the baseline) than background fluorescence (F0). Consequently, the greater the quantity of target DNA in the starting material, the faster a significant increase in fluorescence signal will appear, yielding a lower Ct (Heid et al., 1996). There are many benefits of using real-time PCR over other methods to quantify target DNA. It can produce quantitative data with an accurate dynamic range of 7 to 8 log orders of magnitude and does not require post-amplification manipulations. To date, real-time PCR is widely used in research and clinical practice (Kaltenboeck & Wang, 2005; Espy et al., 2006; Schuler & Dolken, 2006).

If real-time PCR is properly designed, the efficiency coefficient is close to the theoretical value of 1 (Livak & Schmittgen, 2001), indicating that errors due to the PCR amplification process are very low. The most common causes of error are the DNA extraction procedures, a key step for real-time PCR, because any error will be amplified exponentially during the PCR process. DNA extraction efficiency varied from kit to kit and test to test, and the extraction process not only was time consuming but also tended to increase the risk of carry-over contamination. Direct PCR without DNA extraction for qualitative use has been reported in several studies (Castley et al., 2005; Ieamkhang et al., 2005). In the present study, our aim was to quantify human hepatitis B virus (HBV) DNA by direct real-time PCR from serum without a DNA extraction procedure. Since haemolysed, lipaemic and icteric serum samples from patients are inevitable in clinical practice, we evaluated the interference of these endogenous substances (also known as the serum index), including conjugated bilirubin, haemoglobin and triglycerides (Powers et al., 1986), in real-time PCR.

Abbreviations: Cp, crossing point; Ct, cycle threshold; CV, coefficient of variation; Fp, fluorescence at the plateau phase; F0, background fluorescence.
METHODS

Blood samples and standards. The present study was performed with the consent of an ethics committee, as well as with the patient’s consent. Serum samples were collected from 91 patients who were HBsAg+ (hepatitis B surface antigen positive), and 23 healthy blood donors who were HBsAg- and anti-HBc- (anti-hepatitis B core negative). HBsAg+ patients were randomly included from chronic HBV carriers attending follow-up sessions for serial monitoring of liver status. Blood samples were centrifuged within 4 h to obtain serum fractions, which were then divided into aliquots and kept at −80 °C prior to testing. Serological tests for HBsAg were performed with a commercial microparticle enzyme immunoassay (CI8200; Abbott Laboratories). HBV DNA standards were included in the commercial kit (Roche Diagnostics) with HBV DNA concentrations of 5.0 × 10^4, 5.0 × 10^5, 5.0 × 10^6 and 5.0 × 10^7 copies mL−1.

DNA template preparation. Nucleic acid was extracted with a commercial kit (Roche Diagnostics), following the manufacturer’s instructions. A 100 μl volume of serum was precipitated with 100 μl reagent A, and then centrifuged at 11 336 g for 10 min. The pellet was resuspended with 25 μl reagent B, heated to 100 °C for 10 min and spun again at 11 336 g for 10 min. The supernatant was used as a source of DNA for PCR.

Real-time PCR. Real-time PCR kits were obtained from Roche Diagnostics. PCR was performed using a Roche Diagnostics LightCycler and the fluorescent signal of PCR products after each cycle was monitored continuously with a fluorescent hydrolysis probe (TaqMan), according to the manufacturer’s instructions. Primer sequences used for HBV were 5′-ACTCGGTTGGACCTCTCTC-3′ and 3′-AAGAAAGGAGGCGATGCACGA-5′, and for the TaqMan probe were 5′-TGGATCTGTCTGGCGGTTTTATCAT-3′. An aliquot of 2 μl serum or extracted nucleic acid template was mixed with 18 μl reaction mixture, and then added to the capillary. All capillaries were sealed and briefly centrifuged before amplification in the LightCycler. The PCR protocol consisted of 45 cycles of 30 s denaturation at 95 °C, and 20 s annealing and extension at 60 °C. The temperature transition rate was 20 °C s−1 for all steps. Fluorescence data were acquired once each cycle at the end of the annealing phase with a detection channel setting of F2/F1.

The influence of serum protein on real-time PCR. To evaluate the influence of serum protein on real-time PCR, we treated the standards (since the standard in HBV viral particle form was not available) with serial diluted serum protein from 5.0 to 50 g l−1 and observed how the fluorescence and Cp changed.

Interference study. Since sera from patients contained endogenous substances, we studied their interference in real-time PCR. According to the protocol described in National Committee for Clinical Laboratory Standards guidance document EP7-P, the maximum concentrations of endogenous substances used in the interference test were: 240 μmol conjugated bilirubin l−1, 4.0 g haemoglobin l−1 and 5.0 mmol triglycerides l−1 (Powers et al., 1986). Interference tests were performed in duplicate using HBV DNA standards and serum specimens.

Data analysis. For data analysis, baseline adjustment was carried out in the ‘proportional’ mode and fluorescence curve analyses were carried out in the ‘fit points’ or ‘second derivation maximum’ mode of the LightCycler. Positive results obtained for the real-time PCR assay using fluorescent hydrolysis probe were characterized by a sigmoid curve, showing an initial rapid exponential increase in fluorescence signal followed by a plateau. Negative reactions did not show any increase in fluorescence signal. A standard curve was created automatically with the LightCycler software in each run by plotting the Fc (or Cp) number against the copy numbers of each standard, and quantification of HBV DNA for unknown samples was inferred from the regression line.

Statistics. All statistical analyses were performed with the spss 11.0 software package for Windows (SPSS). Comparison between the innovative method and the classical one was by χ² test. A two-tailed P value less than 0.05 was considered statistically significant.

RESULTS

The dynamic characteristics of direct real-time PCR

In this study, direct real-time PCR refers to the real-time PCR performed on unextracted serum. The direct real-time PCR curve was generally of log or sigmoid shape, with a slightly clockwise shift that was similar to the normal real-time PCR curve (Higuchi et al., 1993) (Fig. 1). It also indicated that the PCR kinetics consist of an early ground phase, exponential growth phase, linear growth phase and plateau phase. Serum matrix raised the F0, and this effect was attenuated gradually along with the cycle increment until positive amplification occurred. Taking an HBsAg+ serum specimen with 73.2 g total protein l−1, for example,
the value of fluorescence at the plateau phase (FP) of direct real-time PCR kinetics was only about 10% of that acquired from normal real-time PCR (Fig. 1). This case is representative of all the clinical specimens, which usually had total protein ranging from 60 to 80 g l\(^{-1}\). Fluorescence values of all negative and some positive reactions were always under the baseline. This made it difficult to do baseline adjustment and crossing line setting, so fit point mode was not suitable for Ct number determination. We used second derivative maximum mode for Cp determination. Neither baseline adjustment nor crossing line setting was required in this method, because the software automatically determined the Cp value by identifying the first turning point of the fluorescence curve.

The influence of serum protein on real-time PCR
Serum protein raised the F0 (Fig. 2) and decreased the FP in a concentration-dependent way (Fig. 3), but the Cp remained unchanged (Table 1).

Interference study
All the tested endogenous substances (conjugated bilirubin, haemoglobin and triglycerides) slightly decreased the initial F0 in a concentration-dependent way. The substances also decreased the FP of direct real-time PCR kinetics, although Cp values remained unchanged (Table 1). The Cp of real-time PCR had a good reproducibility even at a relative
fluorescence as low as 6%. Bilirubin decreased the fluorescence value to an undetectable baseline when it was present in concentrations above 240 μmol l⁻¹.

**Performance characteristics and efficiency**

The performance of the direct real-time PCR method was determined from Cp values obtained with HBV DNA standards and known samples (known samples were diluted tenfold from 10⁷ to 10² copies ml⁻¹). All calibration curves showed linearity over the entire quantification range with a correlation coefficient \( r^2 = 0.99 \), indicating a precise log-linear relationship. The linear correlation between the cycle number (\( y \)) and the input concentration (copies per reaction) of HBV DNA template for classical real-time PCR and direct real-time PCR were

\[
y = -3.504 \log_{10} x + 47.89 \quad (r = -1.00) \quad \text{and} \quad y = -3.569 \log_{10} x + 46.63 \quad (r = -1.00)\]

Calculated according to the literature, the amplification efficiencies were 93.0 and 90.6%, respectively. When known samples were diluted to 2 × 10⁵ copies ml⁻¹, they could still be detected in the presence of serum protein. Pooled positive serum was tested 20 times using both methods, giving intra-assay CVs of 5.4 to 1.6 %, and inter-assay CVs of 6.5 to 2.3 %, respectively, for real-time PCR with (normal/classical) and without (direct) DNA extraction (Table 2).

To determine the clinical performance of direct serum real-time PCR, HBV DNA concentrations were measured in serum samples from 91 HBsAg⁺ patients as well as 23 healthy controls. Results were compared to those determined by classical real-time PCR. Taking 1.0 × 10⁵ copies ml⁻¹ as the positive threshold, the seroprevalence of HBV DNA among HBsAg⁺ patients was 65.6 % (63/91) by both methods. The rates of positive conformability and negative conformability between the two methods were both 100 %. The log-transformed HBV DNA concentrations of 63 samples between the two methods demonstrated a good correlation (Fig. 4; \( r^2 = 0.863; P < 0.001 \)). All 23 healthy controls tested negative by both methods, yielding a specificity rate of 100 %.

**DISCUSSION**

The TaqMan hydrolysis probe, which is labelled at its 5’ end with a fluorochrome, emits fluorescent light upon hydrolysis allowing detection of PCR product accumulation. This sequence-specific probe is labelled with a reporter dye on the 5’ end and a quencher dye on the 3’ end, which allows the quencher to reduce the reporter fluorescence intensity by fluorescence resonance energy transfer when the probe is intact. When annealed to the

**Table 1. Fluorescence and Cp interference by serum endogenous substances**

<table>
<thead>
<tr>
<th>No. of DNA copies ml⁻¹</th>
<th>Serum protein (g l⁻¹)</th>
<th>Haemoglobin (g l⁻¹)</th>
<th>Bilirubin (μmol l⁻¹)</th>
<th>Triglyceride (mmol l⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0</td>
<td>10</td>
<td>50</td>
<td>80</td>
</tr>
<tr>
<td>FP (%)</td>
<td>5 × 10⁴</td>
<td>85.8</td>
<td>58.1</td>
<td>15.3</td>
</tr>
<tr>
<td>FP (%)</td>
<td>5 × 10⁵</td>
<td>94.7</td>
<td>75.6</td>
<td>18.4</td>
</tr>
<tr>
<td>FP (%)</td>
<td>5 × 10⁶</td>
<td>98.0</td>
<td>77.4</td>
<td>20.9</td>
</tr>
<tr>
<td>FP (%)</td>
<td>5 × 10⁷</td>
<td>100</td>
<td>84.7</td>
<td>25.5</td>
</tr>
<tr>
<td>CP (Cp)</td>
<td>5 × 10⁴</td>
<td>30.0</td>
<td>29.8</td>
<td>30.6</td>
</tr>
<tr>
<td>CP (Cp)</td>
<td>5 × 10⁵</td>
<td>27.2</td>
<td>27.3</td>
<td>27.5</td>
</tr>
<tr>
<td>CP (Cp)</td>
<td>5 × 10⁶</td>
<td>24.5</td>
<td>24.5</td>
<td>25.0</td>
</tr>
<tr>
<td>CP (Cp)</td>
<td>5 × 10⁷</td>
<td>20.9</td>
<td>20.8</td>
<td>21.0</td>
</tr>
</tbody>
</table>

To determine the clinical performance of direct serum real-time PCR, HBV DNA concentrations were measured in serum samples from 91 HBsAg⁺ patients as well as 23 healthy controls. Results were compared to those determined by classical real-time PCR. Taking 1.0 × 10⁵ copies ml⁻¹ as the positive threshold, the seroprevalence of HBV DNA among HBsAg⁺ patients was 65.6 % (63/91) by both methods. The rates of positive conformability and negative conformability between the two methods were both 100 %. The log-transformed HBV DNA concentrations of 63 samples between the two methods demonstrated a good correlation (Fig. 4; \( r^2 = 0.863; P < 0.001 \)). All 23 healthy controls tested negative by both methods, yielding a specificity rate of 100 %.

**Table 2. Comparison of the two real-time PCR methods**

<table>
<thead>
<tr>
<th></th>
<th>Real-time PCR</th>
<th>Direct real-time PCR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Log-linear relationship</td>
<td>-3.504</td>
<td>-3.569</td>
</tr>
<tr>
<td>Slope</td>
<td>47.89</td>
<td>46.63</td>
</tr>
<tr>
<td>Intercept</td>
<td>0.93</td>
<td>0.91</td>
</tr>
<tr>
<td>Amplification efficiency</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cp reproducibility (n=20)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intra-assay, mean ± SD [CV (%)]</td>
<td>27.14 ± 1.46 (5.4%)</td>
<td>27.63 ± 0.45 (1.6%)</td>
</tr>
<tr>
<td>Inter-assay, mean ± SD [CV (%)]</td>
<td>27.41 ± 1.78 (6.5%)</td>
<td>27.36 ± 0.64 (2.3%)</td>
</tr>
<tr>
<td>DNA extraction</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Performance time (h)</td>
<td>2.0</td>
<td>1.0</td>
</tr>
</tbody>
</table>
target sequence, bound and quenched TaqMan probe will be degraded by Taq polymerase during the extension step of PCR. Probe degradation allows for separation of the reporter from the quencher dye, resulting in increased fluorescence emission following each PCR cycle (Heid et al., 1996). The mechanism by which HBV DNA in intact viral particles can be quantified may be that the HBV particles in the PCR solution release DNA while they are heated during the PCR procedure. Serum proteins are macro biomolecules and may act in one of two ways to interfere with the fluorescence reading. On one hand, they may bind directly to the probe sequence between the reporter and quencher dye, scattering the light emitted from the reporter dye. On the other hand, some proteins may absorb excitation light from the instrument and emit light at a spectrum overlapping that of the reporter dye, raising the excitation light from the instrument and emit light at a dye. On the other hand, some proteins may absorb directly to the probe sequence between the reporter and quencher dye, resulting in increased fluorescence emission following each PCR cycle (Heid et al., 1996). The mechanism by which HBV DNA in intact viral particles can be quantified may be that the HBV particles in the PCR solution release DNA while they are heated during the PCR procedure. Serum proteins are macro biomolecules and may act in one of two ways to interfere with the fluorescence reading. On one hand, they may bind directly to the probe sequence between the reporter and quencher dye, scattering the light emitted from the reporter dye. On the other hand, some proteins may absorb excitation light from the instrument and emit light at a spectrum overlapping that of the reporter dye, raising the initial F0. With the increment of PCR steps, proteins become denatured at high temperatures and may be inclined to form deposits. Deposit particles scatter the excitation light from the instrument, thus little light is absorbed by the reporter dye, yielding little fluorescence. A reporter dye, such as FITC, has a fluorescence excitation spectrum peak at 493 nm and emission spectrum peak at 525 nm. Bilirubin has an absorption band ranging from 450 to 490 nm, whereas haemoglobin has an absorption band ranging from 510 to 550 nm. So, bilirubin quenches fluorescence by absorbing excitation light from the instrument, whereas haemoglobin quenches fluorescence by absorbing light emission from the reporter dye. Triglycerides may dominantly quench fluorescence by scattering excitation light. However, the exact mechanism of each endogenous substance remains to be further elucidated.

The second derivative maximum method for quantification is automatically determining the Cp for individual samples. This was achieved by a software algorithm that identifies the first turning point of the fluorescence curve, which serves as the Cp in this calculation method. Neither baseline adjustment nor crossing line setting is required in this method. The software copes with effects and artefacts such as spikes in the log-linear phase, and noisy plateaus or curves where the plateau phase has not yet been reached. No Cp value will be displayed for a curve with either a slowly increasing or a slowly decreasing fluorescence or a noisy background, which does not allow a considerable fluorescence value to be attained. The second derivative method is a more appropriate method for direct serum real-time PCR, because serum matrix reduced the emitted fluorescence below the acceptable threshold and, sometimes, the Cp value could not be obtained by fit point mode. Increased amounts of circulating DNA have been found in a variety of disorders including cancer, autoimmune disease and infection (Sozzi et al., 2003). It has been predicted for years that simplification of detection methods may widen the availability of this type of testing (Lo, 2000). Our study has further demonstrated the validity of direct serum real-time PCR.

In the present study, we showed a high level of concordance between direct serum real-time PCR and classical real-time PCR during HBV DNA quantification. We also demonstrated that this method could be performed on the LightCycler instrument. By omitting DNA extraction and gel electrophoresis, this method allows for substantial labour and cost savings. In conclusion, direct serum real-time PCR is a simple, economical, time-efficient, sensitive and accurate method for the quantification of serum HBV DNA.

**REFERENCES**


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**Fig. 4.** The relationship between direct PCR and normal PCR. HBV DNA concentrations were log transformed. The time required to perform the direct serum real-time PCR assay was only about 1 h. Comparing direct serum and normal real-time PCR after DNA extraction, we found that the first method saved 1 h.

\[ y = 0.53 + 0.93 \times x \quad r^2 = 0.86 \]


