Novel assay of competitively differentiated polymerase chain reaction for screening point mutation of hepatitis B virus

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**Abstract**

**AIM:** Point mutation, one of the commonest gene mutations, is the most important molecular pathogenesis of cancer and chronic infection. The commonest methods for detection of point mutation are based on polymerase chain reaction (PCR). These techniques, however, cannot be used in large scale screening since they are neither accurate nor simple. For this reason, this study established a novel method of competitively differentiated PCR (CD-PCR) for screening point mutation in clinical practice.

**METHODS:** Two competitively differentiated primers for mutant-type and wild-type templates respectively with an identically complemented region in 3’ end except for last 2 base pairs and a different non-complemented region in 5’ end were designed. Thus, competitive amplification might be carried out at a lower annealing temperature at first, and then differentiated amplification at a higher annealing temperature when primers could not combine with initial templates. The amplification was performed in one-tube. The products of CD-PCR were detected using microplate hybridization assay. CD-PCR was evaluated by detecting G1896A variant of hepatitis B virus (HBV) in form of recombinant plasmids and in sera from patients with hepatitis B, and compared with allele-specific PCR (AS-PCR) and competitive AS-PCR.

**RESULTS:** CD-PCR was successfully established. It could clearly distinguish wild-type and mutant-type plasmid DNA of G1896A variant when the amount of plasmid DNA was between 10^2-10^4 copies/reaction, while for AS-PCR and competitive AS-PCR, the DNA amount was between 10^2-10^4 copies/reaction. CD-PCR could detect one copy of G1896A variant among 10-100 copies of wild-type plasmid DNA. The specificity of CD-PCR was higher than those of AS-PCR and competitive AS-PCR in the detection of HBV G1896A variant in sera from patients with hepatitis B. CD-PCR was independent of the amount of HBV DNA in serum. HBV G1896A variant was more often found in HBeAg(-) patients with a lower level of detectable viremia than that with a higher level of detectable viremia (P=0.0192).

**CONCLUSION:** CD-PCR is more specific since it is less influenced by the amount of initial templates and the cross amplification between mutant- and wild-type amplified products. It is also simple and time-saving. Thus, CD-PCR might be useful in routine gene typing and point mutation screening. HBV G1896A or other more important mutations have to be routinely detected in patients with a detectable level of viremia after HBeAg/antibody conversion in clinical practice.

**INTRODUCTION**

Point mutation is one of the commonest gene mutations not only in human genome but also in genome of pathogens. It is the most important molecular pathogenesis of cancer and chronic infection\[1-3\]. The commonest method for detecting point mutation is based on polymerase chain reaction (PCR), which can be divided into two types. One type is to analyze the PCR products using direct sequencing, restriction fragment length polymorphism and single strand conformation polymorphism\[4-7\]. This type of techniques is usually accurate, but cannot be used in routine clinical examination since it needs additional manipulation. The other type is allele specific-PCR (AS-PCR), sequence specific-PCR or amplification refractory mutation system\[8-13\]. This type of techniques is usually simple, but its specificity is influenced by the amount of initial templates of PCR. This deficiency was not so obvious when this type of methods was used to detect point mutation or gene typing of human gene in the past since the initial templates in these samples could be controlled\[8-12, 13\]. Some investigators, however, have made efforts to improve AS-PCR by introducing competitive mechanism, and established the so-called bidirectional AS-PCR and competitive PCR in order to decrease the influence of the amount of initial templates on its specificity\[14-19\]. AS-PCR has been used to detect point mutation of pathogens recently\[10, 20-23\]. However, the specificity of AS-PCR in detection of pathogen mutations was challenged by uncontrollable amount of initial templates and the requisition of high sensitivity. For these reasons, a novel method of competitively differentiated polymerase chain reaction (CD-PCR) was established in this study and compared with AS-PCR and competitive AS-PCR in the detection of G1896A variant of HBV.

**MATERIALS AND METHODS**

**Materials**

One hundred serum samples with (hepatitis B surface antigen) HBsAg(+), anti-(hepatitis B e antigen) HBe(+) and anti-(hepatitis B core antigen) HBe(+) and 40 serum samples without HBV serum markers were collected. The serum markers were demonstrated by enzyme-linked immunoabsorbent assay. Recombinant plasmid pG1896A was constructed as described...
before\cite{23}], TZ19U-HBV that contained double copies of HBV DNA (adw) was a gift from Professor Huang Zhimin (Zhongshan University, Guangzhou, China) and was used as wild-type DNA control. T4 DNA ligase and pfu DNA polymerase were purchased from Promega (USA). DNA gel extraction kits and plasmid isolation kits were purchased from Qiagen (Germang). Anti-digoxigenin and anti-fluorescein labeled with horseradish peroxidase were purchased from Roche (USA). Primers shown in Table 1 were designed with the Omega 2 software and synthesized in Biosbio, Biological Engineering Company (Shanghai, China).

Table 1 Primers and probes for detecting of HBV G1896A variant

| Denomination | Sequence (5'→3') |
|--------------|------------------|
| BIO-PCP      | BIO-GAGACTCTCAA GGCTTCTCGA TACAGAAGCTAGG |
| PCA          | GCAGTATGGAAGTG GAAGCAATGCTCAC |
| DIG-PCMd     | DIGCTCAGCTCTAC AGTTGTTGCG TGGCTTCA |
| DIG-PCMc     | DITCCTGCTCTGG TGGCACCTCA |
| FLU-PCWd     | FLU-GTCGCTGTCTGTTGCTGGTGGCTTGGT |
| FLU-PCWc     | FLU-GTGCTCTGG TGGCCTTGG |
| PCS          | CCACC GGCGGCTTACAC |
| PCSc         | CGGGATATCCATCC GAGACGCCCCTACAG |
| PCAc         | CGCCAAAGCTTGACTGATGGTGAGTGATGCAATG |

BIO: the abbreviation of biotin; DIG: the abbreviation of digoxigenin; FLU: the abbreviation of fluorescein.

Methods

Principles of CD-PCR Two competitively differentiated primers (CDP) with different labels in 5' end were designed. CDP had a complemented region of 17-20 bp long with the normal bases of the templates. The annealing temperature of this part was about 52-54 °C. There was also a non-complemented region of 14 bp long, which was different from each other in 5' end. The total annealing temperature of CDP was about 75-80 °C. PCR was performed in one tube. Competitive amplification was performed first when CDP competed for templates under lower annealing temperature. After certain amounts of products of each primer were obtained, differentiated amplification was then performed under higher annealing temperature when CDP could only use its own products as templates. This amplification model would keep the products ratio of mutant and wild templates and allow the templates to be fully amplified.

G1896A variant detected by CD-PCR Wild-type and G1896A mutant-type plasmids were used for the optimization of CD-PCR. A total volume of 30 µl was used in the final conditions of PCR reaction. The reaction mixture consisted of 10 mmol/L Tris-HCl, pH 8.5, 50 mmol/L KCl, 1.5 mmol/L MgCl2, 0.2 mmol/L dNTPs, 2U pfu DNA polymerase, 20 pmol FLU-PCWd, 20 pmol DIG-PCMd, 20 pmol PCA and 5 μl plasmid or extracted DNA. The cycling conditions were as follows: 3 cycles (first set) at 94 °C for 40 s, at 52 °C for 40 s and at 72 °C for 90 s followed by 35 cycles (second set) at 94 °C for 40 s, and at 72 °C for 90 s. The PCR products were then hybridized with solidified biotin-labeled probe PCR in two wells of microplate. The color reaction was obtained after captured PCR products reacted with anti-DIG or anti-FLU respectively, which were conjugated with horseradish peroxidase.

G1896A variant detected by competitive AS-PCR Except for the replacement of primer FLU-PCWd and DIG-PCMd with primer FLU-PCWc and DIG-PCMc, the components of competitive AS-PCR were similar to CD-PCR. The cycling conditions were as follows: 25 cycles (first set) at 94 °C for 40 s, at 60 °C for 40 s, and at 72 °C for 90 s followed by 10 cycles (second set) at 94 °C for 40 s, at 52 °C for 40 s, and at 72 °C for 90 s. The PCR products were then detected similar to CD-PCR. G1896A variant detected by AS-PCR Except for the absence of FLU-PCWc, the components and the cycling conditions were similar to competitive CD-PCR. The PCR products were visualized under an ultraviolet transilluminator.

Quantification of serum HBV DNA by fluorescent quantitative PCR The HBV DNA level of 200 sera was detected using fluorescent quantitative PCR following the instruction (Taqmen, Roche).

DNA sequencing A fragment of HBV precore in positive or negative sera by all three methods and in sera that were positive for AS-PCR and competitive AS-PCR, and negative for CD-PCR was analyzed using DNA sequencing after it was amplified using primers PCSc and PCAc and cloned into plasmid pUC19.

RESULTS

Detection of HBV G1896A variant in recombinant plasmid DNA

The typical results of HBV G1896A variant detected by means of AS-PCR, competitive AS-PCR and CD-PCR are shown in Figure 1. To detect G1896A variant, the sensitivity of AS-PCR was 104 copies/reaction, and there was non-specific amplification when the amount of wild-type initial templates was higher than 105 copies/reaction. For competitive AS-PCR, the wild-type and mutant-type plasmid DNA could be clearly distinguished when the initial templates were 105-106 copies/reaction. CD-PCR could clearly distinguish them in the range of 104-105 copies/reaction. There was some degree of non-specific reaction when the amount of initial templates was 106 copies/reaction, but the difference of color intensities between wild-type and mutant-type product wells was obvious. CD-PCR could detect one variant G1896A copy among 100 wild-type copies of DNA when the initial templates were 104-105 copies/reaction, but one out of 10 copies when the initial templates were 105-106 copies/reaction.

Figure 1 HBV G1896A variant detected by AS-PCR, competitive AS-PCR and CD-PCR using its wild-type or mutant-type recombinant plasmid DNA as objects. CP: common product band; MP: mutant product band; W: wild-type products; M: mutant-type products.

Detection of HBV G1896A variant in sera

Among 200 sera, HBV DNA was positive in 104 cases using AS-PCR, in 119 cases using competitive AS-PCR and in 122 cases using CD-PCR. The sensitivity for HBV DNA detection by CD-PCR was similar to that of competitive AS-PCR, but higher than that of AS-PCR (χ²=5.47, P=0.0193). The results of G1896A variant in sera are shown in Table 2. Among the 104 cases with positive HBV DNA detected by all the three methods, there were 33 positive cases, and 31 negative cases.
of G1896A variant detected by both CD-PCR and AS-PCR, and 40 cases that were positive by AS-PCR but not by CD-PCR.

**Table 2** HBV G1896A variant and its relationship with HBV serum markers

|                  | Case                  | CD-PCR(%) | CAS-PCR(%) | AS-PCR(%) |
|------------------|-----------------------|-----------|------------|-----------|
|                  | HBV G1896A DNA(+) variant | HBV G1896A DNA(+) variant | HBV G1896A DNA(+) variant | HBV G1896A DNA(+) variant |
| HBeAg(+)         | 60                    | 100.0     | 6.7        | 100.0     | 63.3<sup>b</sup> | 93.3 | 55.0<sup>b</sup> |
| HBeAg(−)         | 100                   | 62.0      | 38.0       | 59.0      | 46.0        | 48.0 | 40.0 |
| HBeAg(+) / HBeAg(−) | 40                    | 0.0       | 0.0        | 0.0       | 0.0         | 0.0 | 0.0 |

<sup>a</sup>P < 0.01, compared with group of CD-PCR, χ<sup>2</sup> test, <sup>b</sup>P < 0.05, compared with group of AS-PCR, χ<sup>2</sup> test.

**The results of DNA sequencing**

HBV G1896A mutation was confirmed by DNA sequencing in 3 positive sera by all the three methods, and excluded in 3 negative sera by all the three methods and in 4 sera that were positive by AS-PCR and competitive AS-PCR, but negative by CD-PCR.

**The relationship between G1896A variant and HBV DNA level**

The relationship between G1896A variant detected by AS-PCR, competitive AS-PCR and CD-PCR and HBV DNA level in the sera is shown in Table 3. More G1896A variants were found in patients with positive HBeAg by AS-PCR and competitive AS-CR. These could be false positives. These false-positive findings occurred more often in patients with a higher level of HBV DNA in serum. CD-PCR seemed more specific since only 4 patients were found to be infected with both wild-type HBV and G1896A variants in HBeAg positive cases. By CD-PCR, HBV G1896A variant was more often found in HBeAg(-) patients with a lower level of viremia (P=0.0192).

**Table 3 Relationship between G1896A variant and HBV DNA level in serum**

|                  | HBeAg(+) (n=56) | HBeAg(−) (n=48) |
|------------------|----------------|----------------|
| ≤10<sup>6</sup> copies/ml (n=44) | >10<sup>6</sup> copies/ml (n=22) | ≤10<sup>6</sup> copies/ml (n=28) | >10<sup>6</sup> copies/ml (n=20) |
| AS-PCR           | 2              | 31             | 21            | 19       |
| Competitive      | 2              | 35             | 23            | 20       |
| AS-PCR           | 2              | 2              | 21<sup>a</sup> | 8        |

<sup>a</sup>Compared with HBeAg(−) group with a level higher than 10<sup>6</sup> copies/ml of HBV DNA in serum, Fisher’s exact possibility test, P =0.0192.

**DISCUSSION**

AS-PCR has been considered as a simplest and quickest method for the detection of known point mutations<sup>5,10-19</sup>. False positive results, however, seem inevitable since the amplification of wild-type templates by allele-specific or sequence-specific primers for mutant-type templates can not be completely prevented. Once non-specific products occur, they will be amplified subsequently. Thus the ratio of specific and non-specific products will decrease, and become even smaller as the plateau of PCR reaches at the late stage. It is understandable that false band will occur when a large amount of initial templates or too many cycles are used in AS-PCR. To improve the specificity of AS-PCR, the specificity of primers was designed by introducing penultimate mismatch<sup>24</sup>, and the cycles of specific amplification were reduced by introducing basic amplification<sup>6,25</sup>. The application of competitive mechanism is another important step to improve AS-PCR.

Competitive AS-PCR and bidirectional or tetra-primer AS-PCR are common assays that have a competitive mechanism<sup>6,14-19,25</sup>. Bidirectional or tetra-primer AS-PCR is in fact a competition for PCR system, while competitive AS-PCR is the competition for both templates and PCR system. However, these PCRs can not exclude the non-specific amplification from initial templates in each cycle. For competitive AS-PCR, it is possible to reduce the influence of initial templates, but a new problem will occur, which is that the initial products will become the templates for non-specific amplification in subsequent cycles. This might be the underlying mechanism that the competitive AS-PCR seemed worse than AS-PCR in this study. Therefore, to reduce the influence of the initial templates and products would be a very important measure to improve AS-PCR assay on specificity of amplification.

For CD-PCR, competitive amplification was performed first, and followed by differentiated amplification, which was different from competitive AS-PCR. This protocol would allow allele- or sequence-specific amplification to begin at a low level of templates. Thus, the influence of initial templates could be controlled to a minimal level. After a few cycles, differentiated amplification was carried out under higher annealing temperature at which primers could not combine with initiated templates and the initial products could not be used as templates for non-specific amplification in subsequent cycles under higher annealing temperature either. These might be the mechanisms that CD-PCR had much better performance than that of AS-PCR and competitive AS-PCR in this study.

The molecular weights of wild-type and mutant-type products were the same, therefore, products of CD-PCR could not be distinguished by electrophoresis. It was convenient and time-saving when microplate hybridization assay was used to demonstrate the products in this study. Automatic detection was possible too when CD-PCR was adapted to real-time fluorescent PCR assay as used in AS-PCR or competitive AS-PCR<sup>26,27</sup>. CD-PCR could also be used by combining with other high resolution techniques<sup>28,29</sup>. It could be used in gene typing for organ transplantation or gene polymorphism in addition to the detection of gene point mutations as AS-PCR<sup>30,31</sup>.

Detectable viremia was found in sera of some patients with hepatitis B after HBeAg/antibody conversion<sup>32-34</sup>. This type of viremia resulted in G1896A mutation in 60.4 % out of such patients, especially in patients with a much lower level of HBV DNA. This result is conformable to the fact that the replication ability of G1896A variant has decreased<sup>35</sup>, and suggests that G1896A or other more important mutations have to be detected in patients with a detectable level of viremia after HBeAg/antibody conversion in clinical practice.

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Edited by Yuan HT and Wang XL