Development of Fok-I based nested polymerase chain reaction-restriction fragment length polymorphism analysis for detection of hepatitis B virus X region V5M mutation

Hong Kim, Seok-Hyun Hong, Seoung-Ae Lee, Jeong-Ryeol Gong, Bum-Joon Kim

AIM: To develop a Fok-I nested polymerase chain reaction (PCR)-restriction fragment length polymorphism analysis (PRA) method for the detection of hepatitis B virus X region V5M mutation.

METHODS: Nested PCR was applied into DNAs from 198 chronic patients at 2 different stages [121 patients with hepatocellular carcinoma (HCC) and 77 carrier patients]. To identify V5M mutants, digestion of nested PCR amplicons by the restriction enzyme Fok-I (GGA↓TGN9) was done. For size comparison, the enzyme-treated products were analyzed by electrophoresis on 2.5% agarose gels, stained with ethidium bromide, and visualized on a UV transilluminator.

RESULTS: The assay enabled the identification of 69 patients (sensitivity of 34.8%; 46 HCC patients and 23 carrier patients). Our data also showed that V5M prevalence in HCC patients was significantly higher than in carrier patients (47.8%, 22/46 patients vs 0%, 0/23 patients, P < 0.001), suggesting that HBxAg V5M mutation may play a pivotal role in HCC generation in

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chronic patients with genotype C infections.

CONCLUSION: The Fox-I nested PRA developed in this study is a reliable and cost-effective method to detect HBxAg V5M mutation in chronic patients with genotype C2 infection.

Key words: Hepatitis B virus; X antigen; Polymerase chain reaction-restriction fragment length polymorphism analysis; VSM mutation; Hepatocellular carcinoma

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Core tip: In the present study, we developed a reliable and cost-effective Fox-I nested polymerase chain reaction-restriction fragment length polymorphism analysis (PRA) method for the detection of VSM from chronic patients with genotype C2 infection. In addition, our epidemiological data based on the Fox-I nested PRA method strongly support the previous reports that VSM may play a very pivotal role in hepatocarcinogenesis, at least in chronic patients infected with genotype C2.

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INTRODUCTION

Hepatitis B virus (HBV) infection is a global health problem, and more than 350 million people are chronic carriers of the virus[1]. Korea is a recognized endemic area of HBV infection, and an extraordinary prevalence of genotype C2 was also reported in this area, which is known to be more prone to mutations and related to more severe liver diseases and a lower antiviral response compared with genotype B[2,3]. Furthermore, the high prevalence of basal core promoter (BCP) double mutations and the presence of a distinct immune response against HBV proteins in the Korean population could lead to the generation of distinct HBV variants that are rarely encountered in other areas, resulting in distinct clinical manifestations in Korean chronic patients[4-19].

Despite its small size, the HBV genome contains four partially overlapping open reading frames producing at least seven viral gene products[20]. Among these, the HBV X antigen (HBxAg) has been the focus of much attention in recent years because it is implicated strongly in hepatocarcinogenesis. HBxAg is a 154-amino-acid protein with an N-terminal negative regulatory domain and a C-terminal transactivation domain. The HBxAg is multifunctional and affects gene transcription, signaling pathways, genotoxic stress responses, cell-cycle control, and apoptosis, and it also plays an essential role in viral replication[21-24].

During the natural course of HBV infection, naturally occurring mutations may occur, and these could be HBV variants that affect prognosis[25-27]. Several reports have demonstrated that specific point mutations, deletions or insertions in the HBxAg gene were related to severe forms of liver disease, such as cirrhosis of the liver and/or HCC[17,28,29]. In particular, our previous study based on a direct sequencing protocol had introduced a novel N-terminal mutation type, V5M/L, with a mutation in codon 5 of HBxAg from Korean chronic patients with genotype C infections[12]. This mutation type was found significantly more frequently in HCC patients than in patients in other disease groups. This finding suggests that it may play a pivotal role in HCC generation during the natural course of HBV chronic infection. For its monitoring among chronic patients, particularly with genotype C2 infections, the development of a new molecular diagnostic method for its detection is needed. For this purpose, in the present study, we developed a new nested PCR-restriction fragment length polymorphism analysis (PRA) method that can detect VSM and applied it to DNAs from Korean chronic patients.

MATERIALS AND METHODS

Study subjects

Among the patients visiting the Jeju National University Hospital from March to November and the Seoul National University Hospital from January to December in 2005, hepatitis B patients were selected in this study. The patients with a history of anti-viral therapy, an alcoholic liver disease, or hepatitis C involving liver disorders were excluded from this research. For an analysis of the correlation between VSM mutants and hepatocellular carcinoma (HCC), 121 serum samples from HCC patients and 77 samples from carriers were collected and stored at -80 °C. The basic biochemical tests for the serum samples were performed, and hepatitis B e antigen (HBeAg), anti-HBe, HBV-DNA and alpha-fetoprotein (AFP) assays were performed in the case of hepatitis B surface antigen (HBsAg)-positive samples. The detection of HBsAg, HBeAg, anti-HBs, and anti-HBe were performed by a chemiluminescence immunoassay (Abbott ARCHITECT, Abbott, IL, United States), and the determination of the serum HBV DNA was quantified by using a Versant HBV DNA Assay version 3.0 (bDNA, Siemens, NY, United States). This study was approved by the Institutional Review Board of Seoul National University Hospital (IRB No. 1404-070-572).

Diagnosis of liver disease

A definitive diagnosis of liver disease was made according to the overall findings, including clinical,
biochemical, and radiological data. Among patients having positive HBsAg, the subjects with positive HBeAg, anti-HBe negative, HBV DNA positive, and normal serum transaminase were clinically defined as HBeAg-positive healthy carriers if there was no evidence of chronic liver disease in the radiologic findings[30]. Hepatocellular carcinoma was diagnosed by clinical findings such as more than 400 pg/mL of alpha-fetoprotein (AFP), typical hyper blood vessels upon computed tomography (CT) or biopsy from the liver[21].

**HBV DNA extraction**

Viral HBV DNA was extracted from 200 μL of the serum obtained from 121 HCC and 77 carrier patients using the QIAamp DNA Blood Mini Kit (QIAGEN, Hilden, Germany). Briefly, 20 μL of QIAGEN Protease (or proteinase K) was added into the 1.5 mL micro-centrifuge tube containing 200 μL of serum sample. Then, 200 μL of buffer AL was added into the sample and was mixed by pulse-vortexing for 15 s. To ensure efficient lysis, it is essential that the sample and Buffer AL are mixed thoroughly to yield a homogeneous solution. After the sample was incubated at 56 °C for 10 min, 200 μL of ethanol (96%-100%, DukSan, Seoul, Korea) was loaded into the sample and was mixed again by pulse-vortexing for 15 s. After mixing, the mixture was carefully applied to the QIAamp Mini-spin column and was centrifuged at 8000 rpm for 1 min. Then, 500 μL of buffer AW1 was carefully added into the column, and the column was centrifuged at 8000 rpm for 1 min. Next, 500 μL of buffer AW2 was added into the column and was centrifuged at full speed (14000 rpm) for 3 min. After the collection tube containing the filtrate was discarded, 50 μL of buffer AE was added into the column and was incubated at room temperature for 1 min; it was then centrifuged at 8000 rpm for 1 min. The eluted DNA was stored at -20 °C and used for polymerase chain reaction (PCR) mixtures as the template.

**PCR amplification and direct sequencing analysis of HBx region**

To analyze the mutation patterns in the HBx gene from 69 patients amplified by nested PCR, the PCR product was directly sequenced for analysis. The first round of PCR was carried out using the sense primer X-pro-F1 (5’-CTC TGC CAA GTG TTT GCT GA-3’; GenBank accession number AY641558, positions 1171-1190 nt) and the antisense primer X-pro-R1 (5’-CAA GGC ACA GCT TGG AGG CT-3’; positions 1886-1905 nt), which amplify a HBV X region, while the second round of amplification was performed using the sense primer X-pro-F2 (5’-TTG CTC GCC GCC GGT CTG GA-3’; positions 1295-1314 nt) and the antisense primer X-pro-R2 (5’-TGA ACA GTA GGA CAT GAA CA-3’; antisense positions 1866-1885 nt) (Figure 1A). PCR was initiated using the i-MAX DNA polymerase (iNtRON Bio, Seoul, Korea) in a 20 μL mixture containing 1X PCR buffer containing 2 mmol/L MgCl₂, 2.5 mmol/L dNTP each, 5 pmol primer each, and 5 U of i-MAX II DNA polymerase. The 2 μL of Template DNA extracted from patients’ serum was added into the mixture. The final reaction mixture was subjected to 30 cycles of amplification (60 s at 95 °C, 45 s at 60 °C, and 90 s at 72 °C) followed by a 5 min extension at 72 °C. A 96-well thermocycler (Model 9600 thermocycler, Perkin-Elmer Cetus, Norwalk, United States) was also used. The obtained PCR products were analyzed by electrophoresis on 1% agarose gels, stained with ethidium bromide, and visualized on a UV transilluminator. For direct sequence analysis, an Applied Biosystems model 373A automatic sequencer and a BigDye terminator cycle sequencing kit (Perkin-Elmer Applied Biosystems, Norwalk, United States) were used for the sequencing. For sequencing reactions, 60 ng of PCR-amplified DNA, 5 pmol of reverse primer (X-pro-R2), and 4 μL of BigDye terminator v2.0 100 RR mix (Perkin-Elmer Applied Biosystems, Norwalk, United States) were mixed. Contents were adjusted to a final volume of 10 μL by adding distilled water, and the reaction was run for 30 cycles of 10 s at 96 °C, 5 s at 60 °C, and 4 min at 60 °C. Determined sequences were aligned with the sequences of five HBV references (GenBank accession no. M57663, AB100695, AB074755, AY247032, AY641558, and X02496) using the multiple-alignment algorithm in the MegAlign package (DNASTAR, WS, United States).

**Detection of V5M mutation by Fok-I PCR-restriction analysis**

To identify V5M mutants conveniently from PCR product, the restriction enzyme Fok-I (GGA TGN9↓) was used in this study. Amplification of the targeted X region was performed under the same conditions in the first round of the nested PCR using the same primer set (X-pro-F1 and X-pro-R1). The second round of PCR was conducted using the sense primer X-pro-F2 (5′-TTG CTC GCC GCC GGT CTG GA-3’; positions 1295-1314 nt) and the antisense primer X-pro-R3 (5′-CGT GCA GCC GCC GGT CTG GA-3’; positions 1895-1914 nt), which could amplify the size of the 290-bp product (Figure 1A). For sequencing reactions, 60 ng of PCR-amplified DNA, 5 pmol of reverse primer (X-pro-R2), and 4 μL of BigDye terminator v2.0 100 RR mix (Perkin-Elmer Applied Biosystems, Norwalk, United States) were mixed. Contents were adjusted to a final volume of 10 μL by adding distilled water, and the reaction was run for 30 cycles of 10 s at 96 °C, 5 s at 60 °C, and 4 min at 60 °C. Determined sequences were aligned with the sequences of five HBV references (GenBank accession no. M57663, AB100695, AB074755, AY247032, AY641558, and X02496) using the multiple-alignment algorithm in the MegAlign package (DNASTAR, WS, United States).

**Statistical analysis**

Statistical analysis of the data in this study was conducted using the SPSS version 21.0 software program (Professional Statistic, Chicago, IL), and the results are expressed as percentages. Frequency
tables were analyzed using the $\chi^2$ tests or Fisher’s exact test. Student’s $t$ test was used when the data showed a normal distribution. In all tests, $P$ values < 0.05 were considered to be statistically significant.

**RESULTS**

**Design of a novel Fok-I nested PRA for the detection V5M mutation in HBxAg**

Previously, 3 polymorphisms, V5(GTG), L5(CTG) and M5(ATG), were reported to be found in the 5th codon of HBxAg in Korean chronic patients[12]. The V5(GTG) type is wild type for genotype C2. Although L5(CTG) is rare in genotype C2, it is also found in the wild types of genotype A, B, C1 and D (Figure 1B). We therefore sought to develop the diagnostic method of Fok-I-based nested PRA to detect the M5(ATG) type (hereafter designated V5M), a genuine mutation among 3 types of polymorphisms in the 5th codon of HBxAg in this study. First, for the nested PCR, the total sequences of four HBV genotype C2 strains retrieved from GenBank were aligned using the SeqManTM II software (DNASTAR) (data not shown), and two primer pairs (outer and inner for nested PCR) for the amplification of an HBxAg fragment were designed using Oligo V 6.5 (Molecular Biology Insights), producing a 290-bp nested PCR amplicon (Figure 1A and Figure 2). Second, to enable the simple separation between the V5M type and 2 wild types, V(GTG) and L(CTG), we developed a novel Fok-I (GGATGN9↓) PRA algorithm using MapDraw (version 3.14; DNASTAR, Madison, Wis.). The Fok-I restriction enzyme can recognize the V5M mutation but not the 2 wild types, V(GTG) and L(CTG). It produced 3 types of PRA pattern. In the presence of wild type [V(GTG) or L(CTG)] alone in blood sample DNA, it can produce one band of undigested 290-bp in an agarose gel. In the presence of V5M mutant type alone having the Fok-I recognition site in blood sample DNA, it can produce two completely digested bands, 186-bp and 104-bp, in an agarose gel. Finally, with the coexistence of both wild type and V5M type, it can produce three bands, an undigested one (290-bp) from the wild type and 2 digested bands, 186-bp and 104-bp, from the
Table 1  Sensitivity of nested polymerase chain reaction assay for the amplification of the 572-bp HBV X antigen fragment

| No. of samples (n = 198) | Sensitivity |
|--------------------------|-------------|
| PCR amplification        |             |
| HCC (n = 121)            | 46 (38.0)   |
| Carrier (n = 77)         | 23 (29.9)   |
|                          | 34.8%       |

PCR: Polymerase chain reaction.

Figure 2  Algorithm of Fok-I nested polymerase chain reaction-restriction fragment length polymorphism analysis method for the detection of V5M mutation in the HBV X antigen region used in the present study.

**DISCUSSION**

Previously, we introduced a total of 5 types of mutations in the X gene (V5M/L, P38S, H94Y, I127T/N, and K130M and V131I) that were significantly related to the clinical severity of chronic patients infected with genotype C2 via a molecular epidemiologic study of Korean chronic patients[12]. Of these, the V5M/L type was first introduced by us and has some properties that are distinct from other mutations. First, this is a genuine HCC-specific mutation as its prevalence in HCC patients was significantly higher than even that of cirrhosis patients or in chronic hepatitis or carriers. Recently, the combination of both BCP double mutations and both types of the V5M mutation, V5M and V5L, has also been reported to increase the risk of HCC by 5.34 times compared with wild type, suggesting that V5M with HBV genotype C2 is a risk factor for the development of HCC and can be used to predict the clinical impact of chronic HBV infection[32]. Second, it is more prevalent in HBeAg-negative patients than in HBeAg-positive patients, suggesting that it may be generated from host immune pressure[12]. Third, of note, it has a positive correlation with the BCP
K130M and V131I[12]. Taken together, it may play a very pivotal role in hepatocarcinogenesis during the HBeAg-negative immune active stage in the natural course of genotype C2 HBV infection. The development of molecular diagnosis for the detection of VSM must therefore be accomplished in genotype C2 endemic areas such as South Korea and China.

For the molecular detection of VSM mutants of the HBV X region, we developed a molecular-based approach, Fok-I nested PRA, which allowed for the rapid detection of the VSM mutant type without sequence analysis. There are some noteworthy advantages in our Fok-I nested PRA method. First, the 5-bp consecutive HBV sequences (GGATG) corresponding to the Fok-I (GGATGN9↓) recognition site of the VSM mutant type are highly conserved among the HBV genotype C2 strains having the V5M mutation, as shown in Figure 1B. When we analyzed the HBxAg sequences from 45 independent chronic patients infected with genotype C2 who had been identified as having the V5M mutation by direct sequencing analysis[12], no mutations were found in the 5-bp consecutive HBV sequences (GGATG) (data not shown). This result could guarantee the enhanced specificity of our Fok-I nested PRA method. Actually, our Fok-I PRA method can detect all of the V5M mutations with 100% specificity compared with the direct sequencing protocol (Table 2). Second, this method can also detect mixed type infections consisting of both the V5M mutant and the wild type in addition to being able to detect V5M mutation infections alone and wild type infections alone.

However, some concerns regarding this assay should also be noted. First, this method may underestimate the frequency of the V5M/L mutation because it is blind to the L(CTG) genotype starting from the CTG in the 5th codon of the X region, which is found in wild-

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**Table 2** Evaluation of Fok-I nested polymorphism analysis assay for the detection of V5M mutations from 69 samples amplified by nested PCR by comparing with the results of the direct sequencing method n (%)

| Type of HBxAg 5th codon | Result for Fok-I nested PRA | Result for Direct sequencing |
|-------------------------|-----------------------------|-----------------------------|
|                         | HCC (n = 46) | Carrier (n = 23) | HCC (n = 46) | Carrier (n = 23) |
| Wild type               | 24 (52.2) | 23 (100)        | 24 (52.2) | 23 (100)       |
| V5M                     | 17 (37.0) | 0 (0)           | 17 (37.0) | 0 (0)          |
| Wild type + V5M         | 5 (10.8)  | 0 (0)           | 5 (10.8)  | 0 (0)          |

PRA: Polymorphism analysis; HCC: Hepatocellular carcinoma; HBxAg: HBV X antigen.
type genotype A, B, C1 or D, but very rare in genotype C2 as shown in Figure 1B. However, given that the V5L genotype is the wild type in genotype A, B or D, it seems more likely and reasonable that unlike the VSM type, it may be one of the wild types in genotype C2 rather than the mutation generated by host immune pressure. Second, despite the nested PCR protocol, our assay had a low level of sensitivity (34.8%, 69/198 patients) (Table 1). This low sensitivity may be due to the absence of the target X region rather than a defect of our assay as previously reported[33] because in 129 samples not amplified by our assay, our repeated attempts to amplify a partial X gene also failed, but the nested PCR protocol targeting the partial S gene could amplify most of them (data not shown). Our epidemiological data based on our Fok-I nested PRA method showed that VSM mutations are significantly more prevalent in HCC patients than in carrier stage patients [47.8% (22/46 patients) vs 0% (0/23 patients), P < 0.001], strongly supporting previous reports that this mutation may play a very pivotal role in hepatocarcinogenesis, at least in chronic patients infected with genotype C2[12,21].

In the present study, we developed a reliable and cost-effective Fok-I nested PRA method for the detection of VSM from chronic patients with genotype C2 infection. In addition, our epidemiological data based on the Fok-I nested PRA method strongly support the previous reports that VSM may play a very pivotal role in hepatocarcinogenesis, at least in chronic patients infected with genotype C2.

## COMMENTS

### Background
The HBxAg V5M mutation proved to be associated with liver disease progression in chronic subjects with genotype C2 using our previous molecular epidemiologic study based on direct sequencing protocol. So, for its monitoring among genotype C2 infected chronic patients, the development of a new molecular diagnostic method for its detection is needed.

### Research frontiers
For molecular epidemiologic study for detection of VSM mutation, in this study, we developed the novel nested polymerase chain reaction (PCR)-restriction fragment length polymorphism analysis (PRA) method that can detect VSM and applied it to DNAs from Korean chronic patients.

### Innovations and breakthroughs
The present epidemiological data based on the Fok-I nested PRA method strongly support the previous reports that VSM may play a very pivotal role in hepatocarcinogenesis, at least in chronic patients infected with genotype C2.

### Applications
The FRET-based real-time PCR for detection of the preS1 deletion developed in this study might have potential for early prediction for risk of liver disease progression in chronic subjects.

### Terminology
The nested PRA method that can detect VSM could be used for molecular epidemiologic purposes. It can permit not only the simultaneous identification of coexisting quasispecies of both wild type and variant, but also the direct identification of target mutations from primary specimens such as serum.

## Peer-review
The authors developed a reliable and cost-effective Fok-I nested PRA method for the detection of VSM from chronic patients with genotype C2 infection, which could be effectively used for its screening instead of direct sequencing protocols. Furthermore, they also confirmed that VSM may play a very pivotal role in hepatocarcinogenesis, at least in chronic patients infected with genotype C2.

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