A Double Polymerase Chain Reaction Method for Detecting African Swine Fever and Swine Vesicular Disease Virus

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Abstract

Purpose: To establish a double polymerase chain reaction (PCR) method for the simultaneous detection of African swine fever virus (ASFV) and swine vesicular disease virus (SVDV).

Methods: By using reference sequences of ASFV and SVDV, this study synthesized parts of the genes connected to the 19-T vector which was inserted into competent DH5α cells to establish recombinant plasmids. Two specific primers of ASFV P72 proteins and SVDV genome were designed to amplify the two target genes. Two pairs of primers and two kinds of recombinant plasmids were added to one PCR reaction system to establish a double PCR assay for detection of the two diseases simultaneously. The double PCR conditions were optimized and the sensitivity and specificity of the assay determined.

Results: The reaction was optimal with a final concentration of 0.36 μM for each primer, and a final annealing temperature of 55.5 °C. The lowest target gene copy number for detecting SVDV and ASFV was 7.6 × 10^2 and 1.5 × 10^5 copies/μL, respectively. The assay has a high level of specificity as only the recombinant plasmids of ASFV and SVDV were amplified and control plasmids for three other diseases - porcine circovirus (PCV), pseudorabies virus (PRV), and porcine parvovirus (PPV) - failed amplification.

Conclusion: This study provides a rapid, sensitive and specific double PCR method for the simultaneous detection of ASFV and SVDV.

Keywords: African swine fever, Swine vesicular disease, Polymerase chain reaction, Recombinant plasmids

INTRODUCTION

African swine fever (ASF) is an acute, febrile, highly contractive and infectious swine disease caused by African swine fever virus (ASFV) [1-3]. ASFV is a double-stranded DNA virus with an outer membrane coating that has a complex 20 sided symmetry, and a viral genome which is 170 kb-190 kb in overall length and which is currently the only known arbovirus with DNA as its nucleic acid [4]. The clinical manifestations of ASF are quite similar to those of swine fever [5], which include acute fever, accompanied with systemic bleeding and hemorrhage in all the organs. The clinical symptoms range from acute, sub-acute and chronic, manifesting as high fever, skin cyanosis, respiratory disorders and neural symptoms. Once swine is infected with ASF, both the morbidity and mortality can reach 100 % [6]. ASF is listed as one of the major seven swine diseases by OIE and is a Class 1 disease in the list of animal pathogenic microbes in China [7].
ASFV can encode more than 34 structural proteins, amongst which the P72 protein can serve as a nucleocapsid protein. The P72 protein is a major structural protein of ASFV, which accounts for 32% of its virion protein content and has a stable antigenicity [8,9]. ASFV is one of the few viruses which cannot react to neutralization reactions and is conserved across different strains [10]. Therefore, it is the most common target antigen for detecting ASFV [11,12].

Swine vesicular disease (SVD) is an acute and febrile infectious disease caused by the swine vesicular disease virus (VDV), which is listed as a Class A infectious disease by OIE [13]. Belonging to small RNA Circoviride Enterovirus, SVDV is a single-stranded positive RNA virus whose genome is 7400 bp in length [14,15]. The genome of SVDV has a large open reading frame which encodes a strain of polyprotein containing 2185 amino acids which produces P1, P2 and P3 by primary pyrolysis. G-C pairs present at low frequency in the base composition, which conforms to that of other enterovirus. The clinical symptoms of this disease feature as blisters or festering of the oral and nasal mucosa and at hoofs causing serious public health problems.

At present, the major laboratory methods for detecting SVD include ELISA detection [16,17], conventional RT-PCR detection and real-time fluorescence RT-PCR detection; each method has associated advantages and disadvantages.

The aim of the paper was to establish a PCR-based assay that can simultaneously detect ASFV and SVDV with high sensitivity and specificity and provide an effective tool for the diagnosis and effective prevention of African swine fever and swine vesicular disease.

### EXPERIMENTAL

#### Materials

2× Taq PCR MasterMix, DNA Marker DL2000, pMD® 19-T Vectors were purchased from Takara (Japan). Agarose was obtained from Biowest, and Column DNA BACK obtained from the Beijing Tian En Ze Gene Technology Co., Ltd. Competent E. coli DH5α was prepared and preserved by Animal Quarantine Laboratory of Sichuan Agricultural University.

#### Design and synthesis of primers

Amplification primers: Primers were carefully designed using DNAMAN according to the sequences data of the ASFV and SVDV published in GenBank (ASFV log-in ID: AY578708 and SVDV log-in ID: EU151461.1). All primers were prepared by TaKaRa. Table 1 shows the sequences of each of the amplification primers.

Synthetic primers of target genes: ASFV (log-in ID: AY578708) and SVDV (log-in ID: EU151461.1) sequences were downloaded from NCBI and primers designed by DNASTAR. The length of the ASFV target fragment was 235 bp, and the length of SVDV target fragment was 358 bp. Table 2 shows the sequences of the synthetic primers of ASFV, and Table 3 shows the sequences of the synthetic primers of SVDV.

#### Synthesis of target segment

The conserved segment sequence fragments of ASFV and SVDV were synthesized in vitro by an overlap extension PCR method using synthetic primers. The primary extension reaction system included 5 μL of 10×Pfu DNA polymerase buffer, 1 μL of Pfu DNA polymerase, 4 μL of dNTP (10 mM), 1 μL of bypass primers (F1-4/R1-4) (10 mM), and 38 μL of double distilled water.

<table>
<thead>
<tr>
<th>Pathogen</th>
<th>Primer sequence</th>
<th>Product size (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ASFV</td>
<td>AP1: 5’-AACAGCAGGTAAACAAGA-3’</td>
<td>235</td>
</tr>
<tr>
<td></td>
<td>AP2: 5’-ATCCTTTTGGCAGATGC-3’</td>
<td></td>
</tr>
<tr>
<td>SVDV</td>
<td>SP1: 5’-CCTCAATTCTCGTGTCG-3’</td>
<td>358</td>
</tr>
<tr>
<td></td>
<td>SP2: 5’-GAACTTGGATGCGTTACACT-3’</td>
<td></td>
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</table>

<table>
<thead>
<tr>
<th>Pathogen</th>
<th>Primer sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>AF1</td>
<td>AACACGCAGGTAAACAAGATATTACTCCCTATTACGGACGCAAGCTGATCTGGGACATAGAGCA</td>
</tr>
<tr>
<td>AF2</td>
<td>CAAACCCCTAAACTACTACCCAGCGGCGGCTCGGATTAAAGCTGCGGGTTGTT</td>
</tr>
<tr>
<td>AF3</td>
<td>CGGTATCCATTCCCTCGGCGGCTGTTTATCACCATAAAAGCTGGATGCGAAGAAAGAT</td>
</tr>
<tr>
<td>AR1</td>
<td>GTATTAGGTGTTAGGTAACGCTCCATTACAGCAGTAAATGAACATTACGTCTTAATGCTCCAGAT</td>
</tr>
<tr>
<td>AR2</td>
<td>GCTGCCTTTTGGTTAACGAGAACGTGAACCTTGGCTATCCCTCCGTATCATTCCCT</td>
</tr>
</tbody>
</table>
Table 3: Synthetic primers of SVDV target genes

<table>
<thead>
<tr>
<th></th>
<th>Sequences</th>
</tr>
</thead>
<tbody>
<tr>
<td>SF1</td>
<td>CCTCAATTCTCCGTCAGCAGAGGAGTGCGGCTATAGTGACAGAGTTAGATCCATCACCT</td>
</tr>
<tr>
<td>SF2</td>
<td>TGTGCAAACGTAGTGGTTGGATATGGTGTGTGGCCAGCTTACCTGAAAGACGAGGAGGCT</td>
</tr>
<tr>
<td>SF3</td>
<td>ACGTGGCCACGTGCAGGTTTTACACACTCGAATCTGTGATGTGGCAGCAAGGCTCGCA</td>
</tr>
<tr>
<td>SF4</td>
<td>GTCAAACATGGGGCTATTCGGGCAGAACATGCAGTACCACTACCTCGGGAGAGCCGGGT</td>
</tr>
<tr>
<td>SR1</td>
<td>TGCACGTGGCCACGTCCGGTTGGGTGGGTTGATCCTCTGCTGTTGCCTCCTCGTCTTTTC</td>
</tr>
<tr>
<td>SR2</td>
<td>TGCACGTGGCCACGTCCGGTTGGGTGGGTTGATCCTCTGCTGTTGCCTCCTCGTCTTTTC</td>
</tr>
<tr>
<td>SR3</td>
<td>AGCCCCATGTTTGACAACGCGTCGGGGAACTTCCACCACCAGCCTGGCGAGCCTTGCG</td>
</tr>
<tr>
<td>SR4</td>
<td>GAACCTGTGATCGCTTACTGCAGTOATATGTGTGTTACCCCGGCTTCCCGGA</td>
</tr>
</tbody>
</table>

All the primers were diluted to a concentration of 10 μM according to the manufacturer’s instructions and stored at -20°C.

The secondary extension reaction system included 20 μL of two adjacent reaction product from the previous bypass and extension reactions each, 1 μL of Pfu DNA polymerase, 1 μL of 10× Pfu DNA polymerase buffer, and 8 μL of dNTP (10 mM).

Primary and secondary extension reactions were conducted at 94°C for 30 s and 72°C for 15 min, respectively.

Amplification and identification of target genes

The 50 μL reaction system contained 25 μL of 2× Taq PCR MasterMix, 18 μL of double distilled water, 1 μL of ASFVP1, 1 μL of ASFVP2, 1 μL of SVDVP1, 1 μL of SVDVP2 and 3 μL of products from the secondary extension reaction. The amplification reaction was performed at 94°C for 4 min (initial denaturation), 94°C for 50 s (denaturation), 55°C for 45 s (annealing), 72°C for 1 min (extension) for 30 cycles ending at 72°C for 10 min (overall extension). PRC products were stored at -20°C.

The PCR products were detected using 1% agarose gel electrophoresis and connecting the products to a pMD® 19-T Vector after gel extraction and purification. A 10 μL reaction system contained 5 μL of solution I, 1 μL of pMD® 19-T Vector, and 4 μL gel extraction products. The reaction system was maintained at 16°C overnight and inserted into competent DH5α cells for cloning. The bacterial liquid was aliquoted into different EP tubes at a ratio of 7:3 (bacterial liquid: glycerine). Samples were processed for sequencing analysis and others were stored at -70°C.

Establishment of double PCR method

An aliquot of 20 μL of the double PCR system contained ASFVP1, ASFVP2, SVDVP1 and SVDVP2 in the same system (the concentration range of the primers is 0.04 - 0.6 μM), 10 μL of 2× Taq PCR MasterMix, 2 μL of ASFV bacterial liquids, 2 μL of SVDV bacterial liquids and double distilled water. In order to optimize the annealing temperature, 50, 51, 52.9, 55.5, 59.1, 62, 63.8 and 65°C were selected to test reactions. The reaction procedure was as follow: 95°C for 5 min, 94°C for 30 s, annealing temperatures for 45 s, 72°C for 30 s, 30 cycles, ending at 72°C for 10 min. The PCR products were detected by 1% agarose gel electrophoresis.

Specificity experiment

The specificity of double PCR with the two pairs of specific primers was assessed by comparing the recombinant plasmids containing the target ASFV and SVDV fragments to the Porcine circovirus (PCV), Porcine pseudorabies (PRV), and porcine parvovirus (PPV).

Sensitivity experiment

To assess the sensitivity, the recombinant plasmids constructs were measured using a ND-1000 ultraviolet spectrophotometer (Nano Drop Co, Ltd, USA) and amplified in a 10-fold serial dilution by PCR using the optimum reaction system and conditions as described above. Quantitative results showed that the concentration of the ASFV and SVDV gel extraction products were 40 ng/μL and 30 ng/μL respectively. The calculation formula of gene copy number concentration is: \( \frac{(6.02 \times 10^{23}) \times \text{(sample concentration ng/μL} \times 10^{-9})}{(\text{DNA length} \times 660)} = \text{copies/μL} \).

RESULTS

Amplification and identification of target genes

The recombinant plasmids of ASFV and SVDV were used as the templates and amplified by the specific primers of ASFV and SVDV. The reaction products were detected by 1% agarose gel electrophoresis. The results showed that ASFV and SVDV amplify fragments at the lengths of 235 bp and 358 bp respectively, which...
conformed to the length of expected target gene fragments (Fig. 1).

![Fig 1: Identification of recombinant plasmids. Note: 1 = negative control; 2 = recombinant plasmids; and 3 = DNA marker DL2000](image)

The previously selected target genes of ASFV and SVDV were compared for homology (in www.NCBI.com) and the results showed 100% homology, indicating that the target genes are correctly synthesized.

**Optimization of double PCR reaction conditions**

**Optimization of annealing temperature**

The optimal anneal temperature results are shown in Fig. 2. In consideration of the effect of temperature on annealing and specificity, the optimum annealing temperature was 55.5 °C.

![Fig 2: Optimization of annealing temperature. Note: M = DNA marker DL2000; 1 = negative control; 2 – 9 = annealing temperature of 65, 63.8, 62, 59.1, 55.5, 52.9, 51 and 50 °C, respectively](image)

**Optimization results of primer concentration**

When the final concentration of each specific primer reached 0.04–0.6 μM, the amplification results are as shown in Fig. 3. For reaction efficiency and reduction of primer dimer formation, the optimal final concentration of primers was shown to be 0.36 μM.

![Fig 3: Optimization of primer concentration. Note: M = DNA marker DL2000; 1 – 12 = respective primer concentration 0.6, 0.48, 0.36, 0.3, 0.24, 0.2, 0.18, 0.16, 0.12, 0.08, 0.06 and 0.04 μM](image)

**Specificity results from double PCR**

PCR amplification of ASFV, SVDV, PCV, PRV and PPV using the optimized double PCR conditions are shown in Fig. 4. Lane 4 demonstrates the only amplification specific products at lengths of 235 bp and 358 bp. The amplification results of PCV, PRV and PPV pathogens are all negative, indicating that the double PCR method is highly specific.

![Fig 4: Specificity results. Note: 1 – 3 = PCV, PRV, and PPV, respectively; M = DNA marker DL2000; 4 = mixed bacterial liquids of ASFV and SVDV](image)
Sensibility experiment on double PCR

The results of sensibility experiment are shown in Fig. 5. It is observed from Fig. 5 that SVDV in lane 2 still presents obvious specific amplification with a sensitivity to the target genes of $7.6 \times 10^2$ copies/µL. ASFV in lane 4 presents obvious specific amplification with a sensitivity to the target genes of $1.5 \times 10^5$ copies/µL.

**Fig 5:** Sensibility results. *Note:* 1 - 10 = ASFV target genes copy numbers, viz, $1.5 \times 10^2$, $1.5 \times 10^3$, $1.5 \times 10^4$, $1.5 \times 10^5$, $1.5 \times 10^6$, $1.5 \times 10^7$, $1.5 \times 10^8$, $1.5 \times 10^9$ and $1.5 \times 10^{10}$ copies/µL, respectively; 1 - 10 = SVDV target genes copy numbers, viz, $7.6 \times 10^1$, $7.6 \times 10^2$, $7.6 \times 10^3$, $7.6 \times 10^4$, $7.6 \times 10^5$, $7.6 \times 10^6$, $7.6 \times 10^7$, $7.6 \times 10^8$, $7.6 \times 10^9$ and $7.6 \times 10^{10}$ copies/µL, respectively; M = DNA marker DL2000

**DISCUSSION**

ASF is a serious disease that can cause huge economic losses in the swine industry. The number of swine in China accounts for half of the total number in the world and our consumption of pork is ever increasing, which requires increasing quantity of pork imported from abroad. Therefore, the development of a healthy and robust swine industry is extremely important and has a significantly influences the food safety and national security in China.

Currently, there is no ASFV in China and ASF epidemics in the adjacent areas between Russia and China. However, the constant spreading of ASF epidemics within Russia and the frequent, rapid and long-distance trading of animals and commodities around the globalized world make it increasingly difficult to guarantee that food is free from pathogenic bacteria, virus or epidemics, such as ASFV. Therefore, we must remain on high alert and conduct risk analysis to prevent ASF from entering in China [18]. Besides, SVD's damage to swine industry has been continuously increased in recent years. Therefore, it is significant for ensuring the healthy development of swine industry in China by establishing a rapid and accurate method for simultaneously detecting ASF and SVD.

To our knowledge, many available techniques such as RT-PCR, real-time RT-PCR and ELISA have been developed for ASFV detection in the endemic areas, but there was few studies on ASFV including the ASFV detecting methods in China and some other non-endemic countries because ASF have not happened in the boundary regions and there is no ASF strains used for researches in China.

This study designs two kinds of specific primers for ASFV P72 proteins and SVDV genome respectively and synthesizes target genes. By optimizing the annealing temperature and other reaction conditions for the double PCR assay, this study guarantees the amplification efficiency of the two couples of specific primers in the specific reaction. The double PCR method established in this study can rapidly detect and identify ASFV and SVDV in one reaction system. It shows good specificity and sensitivity, and can be directly used for the detection of clinical epidemic materials in samples, offering an effective tool for the diagnosis, and prevention and control of ASF and SVD. It was the first time to establish an assay to detect the two diseases simultaneously in China. In conclusion, the designed primers and the double PCR described in this paper supplied important materials and a rapid useful technological reserve for ASFV and SVDV detection and set basis for the studies about ASFV detection in China.

**CONCLUSION**

The findings indicate that double PCR approach is a rapid, sensitive and specific method for detecting ASFV and SVDV simultaneously.

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