Generation of Synthetic Peptide-Specific Antibody for the Development of A Southern Rice Black-Streaked Dwarf Virus Diagnostic Test

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Abstract

Southern rice black-streaked dwarf virus (SRBSDV) causes severe epidemiological disease on rice with the infected area up to millions of hectares in South China and North and Central of Vietnam. So far, there are no effective, cheap, quick, and practicable methods for diagnosing SRBSDV. The conventional RT-PCR technique is the most popular method for detecting SRBSDV with high accuracy. However, it is hard to apply this method for large-scale SRBSDV diagnosis because of the requirements of expensive reagents and instruments, as well as complex procedures. Meanwhile, SRBSDV diagnostic techniques based on antigen detection have outstanding advantages due to their low cost, easy manipulation, and wide application possibility. Today, there are still no commercially available specific antibodies to SRBSDV. In a previous study, to develop the SRBSDV diagnostic technique by the ELISA technique, a SRBSDV specific antibody was generated by a recombinant P10 envelope protein (66kDa), which has a titer of 1:5,000. In this study, we continued to study the production of SRBSDV specific polyclonal antibodies from small antigen–rich peptides from the SRBSDV P10 envelope protein. The resulting purified antibody can specifically bind to the P10 protein and at the diluted concentration of 1:100,000 it can detect SRBSDV in infected rice samples via the dot-blot technique. Our research results open up new opportunities for proactive antibodies to develop a SRBSDV membrane rapid diagnostic kit.

Keywords

Polyclonal antibody, rice, P10 envelope protein, SRBSDV

Introduction

Southern rice black-streaked dwarf virus (SRBSDV), a member of the genus *Fijivirus* in the family Reoviridae, is one of the most...
dangerous harmful rice viruses and is distributed mainly in southern Asia including China, Japan, and Vietnam (Guo et al., 2008; Zhang et al., 2008; Ngo Vinh Vien et al., 2009; Cuong et al., 2010; Wang et al., 2010; Matshukura et al., 2013). SRBSDV was first observed in Vietnam in 2009 and caused severe epidemic disease in huge rice-growing areas in North and Central Vietnam. So far, the development of an SRBSDV-resistant rice variety is not yet available; therefore, disease management is still primarily aimed at controlling the source of the pathogen, especially the disease vector-like white-backed planthopper (Sogatella furcifera).

The genome of SRBSDV is about 29,124bp in length, consisting of 10 double-stranded RNA segments (from S1 to S10), ranging in size from 1.8 to 4.5kb. In particular, the segment S10 has a length of 1,797bp, containing 1,674 nucleotides and an ORF encoding proteins that form the outer shell of the virus molecule with a corresponding molecular weight of 62.6kDa. The P10 protein is highly conservative, and has been identified as having specific interactions with the vectors of SRBSDV; therefore, it plays an important role in the evolution of the virus. For this reason, the P10 protein is often chosen as the target for developing SRBSDV diagnostic methods (Yin et al., 2011).

Currently, the most common SRBSDV diagnostic technique is reverse transcription-polymerase chain reaction (RT-PCR) using specific primers designed based on a highly conservative sequence on the S10 segment of the viral genome (Zhang et al., 2008; Ngo Vinh Vien et al., 2009; Cuong et al., 2010; Anh et al., 2011). SRBSDV diagnostic techniques using real-time RT-PCR are also being applied to quantify viruses for early diagnosis (Zhang et al., 2013). Chinese laboratories have produced anti-SRBSDV antibodies using several synthetic peptide fragments of the P10 protein and developed a diagnostic technique using dot-ELISA (Wang et al., 2012).

In previous studies, we have successfully created polyclonal antibodies against SRBSDV from the recombinant P10 envelope protein segment to detect SRBSDV in ELISA tests (Do Thi Hanh et al., 2015). However, in order to develop an SRBSDV rapid diagnostic kit, the antibodies produced need to be specific and have a high titer to ensure the accuracy and sensitivity of the diagnostic kit. In this research, we continued to generate the polyclonal antibodies against SRBSDV based on small conservative antigen-rich peptides derived from the P10 protein of representative Vietnam SRBSDV isolates. The resulting antibody will be the premise for developing an SRBSDV rapid diagnostic kit in future.

**Materials and Methods**

**Materials**

The SRBSDV artificially infected rice samples, with and without visible symptoms that showed positive results in RT-PCR tests, were supported by the Research Center for Tropical Plant Pathology, Vietnam National University of Agriculture. Laboratory 8-week-old mice were ordered from the National Institute of Hygiene and Epidemiology. The recombinant peptide was synthesized by the GeneCreate Company (China).

**Methods**

**Preparation of bacterium and rice extracts**

Small-scale extractions were conducted to test for dot blotting and western blotting experiments. For the rice extraction, approximately 1.0 gram of fresh rice leaf tissue was ground into a powder in liquid nitrogen and homogenized with 1.0mL of 1X PBS buffer. For the bacterium extraction, the recombinant E. coli cell lines containing either the pET28a or pET28a/P10 vector were cultured for protein expression as previously described (Pham Thanh Tam et al., 2013), and then lysated in 1X PBS buffer by using a sonicator device. All mixtures were then centrifuged at 16,000 × g for 10min and the supernatant was kept as a testing sample.

**Design of the antigenic peptide fragment**

The conservative amino acid sequence on the P10 protein of SRBSDV was identified by BioEdit 2.0 software based on the published S10 segment sequences of the Vietnam SRBSDV isolates (Nguyen Hoang Quang et al., 2013). The
hydrophilic property and surface probability of the amino acids on the P10 protein molecule were analyzed by the Antibody Epitope Prediction tool (http://tools.immuneepitope.org/). The antigenicity of the P10 protein molecule was analyzed by the Antibody Epitope Prediction tool (http://tools.immuneepitope.org/) and the Predicting Antigenic Peptides tool. The antigen epitope was predicted by the Antibody Epitope Prediction tool (http://tools.immuneepitope.org/) and the BepiPred-2.0 tool (http://www.cbs.dtu.dk/).

Immunization of mice with the synthetic peptide

Immunization experiments were conducted as previously described (Amero et al., 1994). Mice were immunized with 20.0µg of purified fused peptide mixed with 200.0µL of PBS buffer (50mM potassium phosphate, 150 mM NaCl, pH 7.25) and an equal volume of Freund's Complete/Incomplete Adjuvant. The mixture was injected into mice at the peritoneal abdomen. Three immunizations were delivered intraperitoneally, administered one week apart. Tail bleeds were performed to check for immunization responses before primary immunization and one week after each re-immunization with a dot blotting assay (Sambrook & Russell, 2001). When a mouse achieved the highest immune response, the entire mouse blood was collected to extract the serum.

Purification of the IgG antibody

The IgG antibodies in the mouse serum were purified by a protein A-sepharose affinity chromatography column (Thermo Fisher Scientific) as described by Amero et al. (1994). The mouse antisera were titrated to pH 8.0 by adding 0.1 volumes of 1 M Tris-HCl (pH 8.0). The column was equilibrated with 10.0mL of 100 mM Tris-HCl buffer (pH 8.0). After loading the sample, the column was washed, respectively, with 100 mM and 10 mM Tris-HCl buffer. The target protein was eluted by 100 mM Glycin-HCl buffer (pH 3.0). The eluted solution was neutralized to pH 8.0 by adding 0.1 volumes of 1 M Tris-HCl buffer (pH 8.0). The purity of the IgG antibody was confirmed by SDS-PAGE gel electrophoresis and western blotting analysis (Sambrook & Russell, 2001).

Western blotting and dot blotting assays

Recombinant proteins in purified and unpurified samples were detected via western blotting assays (Sambrook & Russell, 2001). The protein samples, after being separated by the electrophoresis method on an SDS-PAGE gel, were transferred electronically to a PVDF membrane in Tris-glycine buffer with 20% methanol. The membrane was immobilized by gently shaking it (for 30 to 60 minutes) in 1% BSA solution and incubated alternately with primary and secondary antibodies for 1 hour. The targeted protein was colored by incubation in p-nitro blue tetrazolium chloride 5-bromo-4-chloro-3-indolyl phosphate (NBT/BCIP) substrate solution.

For the dot blotting assay, the PVDF membrane was activated by soaking it in methanol for 10 seconds. Solutions containing the target protein (prepared in 1X PBS buffer) were dropped directly onto the membrane. Membrane blocking, antibody probing, and detection steps were similar to the western blotting assay (Sambrook & Russell, 2001).

Results and Discussion

Design of the peptide antigen

The P10 protein of SRBSDV is a common target in diagnostic assays of SRBSDV using specific antibodies (Zhang et al., 2013; Do Thi Hanh et al., 2015). To create antibodies specifically binding to the P10 protein of Vietnamese SRBSDV isolates, the peptide fragment antigen was screened according to whether it was located on the (i) antigenic or (ii) conservative region of the P10 protein. In a previous study, we evaluated the S10 genetic diversity of SRBSDV isolates collected from rice growing in different ecological areas of Vietnam (Nguyen Hoang Quang et al., 2013). Based on those published sequences, the conservative region on the P10 protein of SRBSDV isolated in Vietnam was identified (Figure 1).

The antigenic regions (epitopes) of a protein consist of residues located on the surface of the protein structure that are capable of causing an immune response and thus, determine the quality
Note: Parts of the deductive amino acid sequences from the P10 protein of ten Vietnamese SRBSDV isolates (Thai Binh 1, Nam Dinh 1, Nghe An, Quang Tri, Hue 1, Son La, Lao Cai, Ninh Binh, Thai Binh 2, and Hue 2) were analyzed by BioEdit 2.0 software. The dots (.) indicate the same amino acid.

Figure 2. The deductive amino acid sequence of the P10 protein.
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Figure 3. Prediction of the antigenicity of the P10 protein

Note: The Hydrophilicity (A) and Surface Accessibility (B) were analyzed by the Antibody Epitope Prediction tool; the red line represents the threshold value. The epitopes and molecular structure of the P10 protein (C) were predicted by the BepiPred 2.0 tool; the letter E represents the position of the epitope. The antigenicity of the P10 protein (D) was analyzed by the Predicting Antigenic Peptides tool; the bar represents the amino acid zone with high antigenicity.

Figure 4. Evaluation of the P10-specific antiserum responses in mice

Note: Pre-immunization (A) and 4-week immunization (B) antisera were diluted at 1:1000. (BL) Blank control (PBS buffer); (SRBSDV): SRBSDV-infected rice extract; (-): Healthy rice extract; (P10) Purified P10 protein.
To confirm the purity of the obtained antibody, the specific binding protein fraction obtained from the A-sepharose protein column (fraction 7) was analyzed by using SDS-PAGE electrophoresis. There were two protein bands shown on the SDS-PAGE gel with the sizes of about 25 and 53kDa (Figure 4B, lane 1), corresponding to the sizes of the light chain and the heavy chain, respectively, of the IgG antibody. This result proved that the IgG antibody had been purified successfully from mouse serum using the A-sepharose protein affinity chromatography system.

Titer and specificity evaluation of the antibody

In order to determine the titer of the archived antibody, the dot blotting test was conducted with both the SRBSDV-infected rice extraction and the purified recombinant P10 protein sample, using a range dilution of the IgG antibody (from 1:10,000 to 1:100,000). The results of the dot blotting test showed that the purified IgG antibody could detect the target protein at the 1:100,000 dilution and 1:80,000 dilution in the purified protein sample and SRBSDV-infected rice extract sample, respectively (Figure 5, column "P10" and "SRBSDV").

To assess the specificity of the polyclonal IgG antibody, a western blotting assay was conducted on the 1:10,000 dilution of the purified antibody, using a total lysate of the P10 expression in *E. coli* Rosetta cells (Pham Thanh Tam *et al.*, 2013) as a testing sample. The results of the western blotting analysis showed that our antibody could identify a single approximately 66 kDa band in the *E. coli* extract sample (Figure 6B, lane 1), as well as in the purified P10 protein sample (Figure 6B, lane 3). In contrast, there was no protein band in the negative control experiment using the crude extract of *E. coli* Rossetta cells carrying empty pET28a vectors (Figure 6B, lane 2). These results proved that our purified IgG antibodies could specifically recognize the recombinant P10 protein of SRBSDV.

To confirm the potential of antibody application in SRBSDV diagnosis, purified IgG antibody (diluted at 1:10,000) was subjected to a dot blotting test against the different rice extracts containing SRBSDV. Colored dots were clearly observed for all the SRBSDV-infected rice samples that showed obvious symptoms (Figure 7, S1 – S4). In contrast, the healthy sample did not exhibit a colored spot (Figure 7, uninfected). In the case of the SRBSDV-infected rice samples without apparent symptoms (Figure 7, NS1 – NS3), they showed lighter spots. These results indicated that the polyclonal anti-P10[15-31] peptide antibody could be used to diagnose SRBSDV.
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Figure 6. Titer of the purified polyclonal IgG antibody against the P10 protein

Figure 7. Specificity evaluation of the purified IgG antibody

Figure 8. Diagnosis of SRBSDV infected rice samples using purified IgG antibody
So far, only Wang et al. (2012) have been successful in the production of specific anti-SRBSDV antibodies by using three peptides with lengths of 20 amino acids, respectively, located at 20-39, 140-159, and 319-339 on the sequence of the P10 protein. The obtained antibody had a titer of 1:500,000 in the ELISA assays. In a previous study, we also used the full-length P10 protein of SRBSDV for mouse immunization and archived IgG antibodies with a titer of 1:6,000. In this study, the anti-SRBSDV antibody was created by a synthetic peptide fragment located at the 15-31 position on the P10 protein, which showed a titer of 1:100,000 in the dot blotting assays. Our antibody detected the presence of SRBSDV in both non-symptomatic and symptomatic rice samples. The archived antibody had a lower titer than that mentioned in the publication of Wang et al., but this could be due to the different titer evaluation methods. Wang et al. (2012) used an ELISA assay for tittering the antibody, in which the measured results were based on the OD_{405nm} value. It is likely that the presence of chlorophyll in the rice samples affected the ELISA results. Therefore, they used the antibody at a dilution of 1:1,500 for the dot blotting diagnosis, and this antibody has not yet been commercialized. Our research results are a significant step in the development of a rapid diagnostic test kit for SRBSDV detection in Vietnam.

Conclusions

A P10[15-31] peptide fragment with a length of 16 amino acids was identified as having high antigenicity on the envelope of the P10 protein of SRBSDV. Polyclonal IgG antibodies against P10[15-31] were successfully generated from laboratory mice and purified via protein A-sepharose column chromatography. The purified antibody had a titer of 1:100,000, and could specifically recognize the recombinant P10 protein in bacterial cell extracts and in SRBSDV in both symptomatic and non-symptomatic SRBSDV-infected rice samples by using dot-blot at the dilution of 1:10,000.

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