Characterization of Two Immunodominant Antigenic Peptides in NSP2 of PRRSV-2 and Generation of a Marker PRRSV Strain Based on the Peptides

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Abstract

Porcine reproductive and respiratory syndrome (PRRS) is a widespread disease with great economic importance in the pig industry. Although vaccines against the PRRS virus (PRRSV) have been employed for more than 20 years, differentiating infected from vaccinated animal remains challenging. In this study, all 907 non-structural protein 2 (NSP2) full-length sequences of PRRSV-2 available from GenBank were aligned. Two peptides, at positions 562–627 (m1B) and 749–813 (m2B) of NSP2, were selected, and their potential for use in differential diagnosis was assessed. Both m1B and m2B were recognized by PRRSV-positive pig serum in peptide-coated enzyme-linked immunosorbent assays. Further epitope identification yielded five overlapping short peptides for the immunodominant regions of m1B and m2B. Using the infectious clone of PRRSV HuN4-F112 as a template, the deletion mutants rHuN4-F112-m1B, rHuN4-F112-m2B, and rHuN4-F112-C5-m1B-m2B were generated and successfully rescued in Marc-145 cells. Growth kinetics revealed that deletion of m1B and m2B did not significantly affect virus replication. Hence, m1B and m2B show potential as molecular markers for developing a PRRSV vaccine.

Key Points

- NSP2-specific peptides m1B and m2B react with PRRSV-positive pig serum.
- Deletion of m1B and m2B has no significant effect on virus replication.
- Peptides m1B and m2B can serve as markers in a diagnostic PRRSV DIVA vaccine.

Introduction

Porcine reproductive and respiratory syndrome (PRRS) is an important viral disease that causes reproductive disorders in sows and respiratory disorders in pigs of all ages (Rossow 1998). Since its discovery in the United States in 1987 and Europe in 1990, the disease has led to huge economic losses in the pig industry (Albina 1997; Benfield et al. 1992; Chueh et al. 1998). Genomic sequencing has identified two PRRS virus (PRRSV) genotypes: PRRSV-1 (European type) and PRRSV-2 (North-American type) (Collins et al. 1992; Nelsen et al. 1999; Wensvoort et al. 1991). PRRSV is a positive single-stranded RNA virus containing a 5′-untranslated region and at least 10 open reading frames (ORFs), including ORF1a, ORF1b, ORF2a, ORF2b, ORF3, ORF4, ORF5, ORF6, ORF7, and ORF5a (Johnson et al. 2011). The polyproteins pp1a and pp1ab, encoded by ORF1a and ORF1b, respectively, undergo hydrolysis to produce 14 non-structural proteins (NSPs): NSP1α, NSP1β, NSP2–NSP6, NSP7α, NSP7β, NSP8–NSP12, and NSP2TF (Fang et al. 2012; Music and Gagnon 2010; Ziebuhr et al. 2000). Although NSP2 is the largest replicase of PRRSV, its gene exhibits significant diversity among different PRRSV strains (Gao et al. 2004). Mutations, insertions, and deletions are common in the middle region of NSP2 (Fang and Snijder 2010; Han et al. 2007).

The first PRRSV was isolated in mainland China in 1996 (Guo et al. 1996), after which the disease spread to all main pig-rearing provinces in the country. In 2006, a more serious and highly pathogenic PRRSV (HP-PRRSV) variant, characterized by a 30-amino acid deletion in NSP2, triggered an outbreak with high
morbidity and mortality among pigs of all ages in some Asian countries (Tian et al. 2007). Subsequent experiments showed that the 30-amino acid deletion in NSP2 was not related to the pathogenicity of HP-PRRSV (Zhou et al. 2009). Recently, NADC30-like PRRSV has emerged as the cause of another epidemic in China (Zhao et al. 2015; Zhou et al. 2015), which has resulted in increased abortions among pregnant sows.

Vaccination is a key strategy for controlling the spread of PRRSV. Approximately ten vaccines against PRRSV have been licensed, including killed vaccines and modified live vaccines (MLV). In general, the latter provide better immune protection than the former but cannot readily distinguish between infected and vaccinated animals. To identify potential positions in NSP2 into which to introduce molecular markers, NSP2 sequences were systematically compared, and two universal regions were selected for antigenic analysis. Importantly, the two regions were deleted without affecting the growth of the PRRSV MLV HuN4-F112 strain in vitro, suggesting that these regions were suitable for developing vaccines capable of differentiating infected from vaccinated animals (DIVA).

**Materials And Methods**

**Cells, virus and antibodies**

Marc-145 cells were grown in Dulbecco's modified Eagle's medium (Invitrogen, Carlsbad, CA, USA) supplemented with 10% fetal bovine serum at 37 °C and 5% CO₂. The modified live vaccine HuN4-F112 strain, the infectious clones of HuN4-F112 and HuN4-F112-C5, were generated in our previous study (Tian et al. 2009; Zhao et al. 2015). Positive pig serum against HP-PRRSV HuN4-F112, NADC30-like PRRSV HeB108 strain, classical swine fever virus, porcine epidemic diarrhea virus, transmissible gastroenteritis virus, porcine circovirus type 2, pseudorabies virus, and porcine parvovirus, as well as a monoclonal antibody (3F7) against PRRSV, were obtained from our laboratory (Wang et al. 2014).

**Dataset and alignment of NSP2 from PRRSV-2**

In our previous study, two amino acid residues at positions 585–586 in the NSP2 of PRRSV HeB108 were deleted during viral passage in Marc-145 cells (unpublished data), indicating that these or nearby residues can be used as markers for next-generation DIVA vaccines. To identify the deletions in circulating PRRSV and MLV strains, all 907 NSP2 full-length sequences of PRRSV-2 available from GenBank in 2020 were downloaded, followed by amino acid sequence alignment and insertion/deletion (indel) analysis using DNASTAR software v7.1.0 (DNASTAR, Madison, WI, USA).

**Immunoreactivity of m1B and m2B peptides**

Indirect enzyme-linked immunosorbent assay (ELISA) was used to evaluate the immunoreactivity of m1B and m2B peptides. All peptides employed in the study were synthesized by GL Biochem, Ltd. (Shanghai, China). ELISA plates were coated with m1B or m2B peptides alone or in combination at a dose of 10 µg/well by incubation in carbonate-bicarbonate buffer (pH 9.6) at 4 °C overnight. The plates were then
blocked with 5% non-fat dry milk in phosphate-buffered saline (PBS) containing 0.05% Tween-20 (PBST) for 1 h at 37 °C. After washing thrice with PBST, 100 μL PRRSV-positive pig serum (1:40 dilution) was added to the wells; the plates were incubated followed by incubation at 37 °C for 1 h, washed again, and then incubated with horseradish peroxidase-conjugated rabbit anti-pig IgG (1:40,000 dilution; Sigma-Aldrich, St. Louis, MO, USA) in PBST at 37 °C for 1 h. Finally, the plates were washed and incubated with 100 μL/well of 3,3',5,5'-tetramethylbenzidine (Invitrogen) for 15 min. The reaction was stopped with 2 M H₂SO₄ (100 μL/well), and the results were read at 450 nm.

**Assessment of specificity**

The specificity of 1B2B-ELISA indirect ELISA was examined using the antisera of the six porcine viruses mentioned above to assess the degree of assay cross-reactivity.

**Immunoreactivity of m1B and m2B truncated peptides**

To identify the immunodominant antigen regions in the m1B and m2B peptides, seven overlapping peptides (m1B1–m1B7) spanning the m1B region were designed. Each peptide was 16 residues long, and the overlapping region between two adjacent peptides spanned eight residues (Table 1). Similarly, seven overlapping peptides (m2B1–m2B7) were designed and synthesized to probe the m2B region. The resulting m1B1–m1B7 and m2B1–m2B7 peptides were used as coating antigens in ELISA, and their reactivity to PRRSV-positive serum was detected as described above.

**Generation of PRRSV m1B or m2B deletion mutants**

Plasmids harboring m1B or m2B deletion mutants were constructed using HuN4-F112 as a template and specifically designed primers (Table 2); a plasmid harboring the m1B and m2B double-deletion mutant was constructed using HuN4-F112-C5 as a template and specifically designed primers. The native m1B and m2B genes in the HuN4-F112 infectious clone were replaced with their deletion variants via overlap PCR amplification using ultra-fidelity DNA polymerase (TaKaRa, Shiga, Japan) as described previously (Yu et al. 2014), followed by digestion with restriction enzymes (Fse I and Nhe I) and ligation. The deletions were confirmed by DNA sequencing. The plasmids were transfected into Marc-145 cells using Xtreme GENE-HP DNA transfection reagent (Roche Applied Science, Basel, Switzerland) as described previously (Zhao et al. 2018).

At 5 days post-transfection, the cells were assayed by indirect immunofluorescence (Gao et al. 2016). Marc-145 cells were incubated with a monoclonal antibody against the M protein of PRRSV and stained with fluorescein isothiocyanate-labeled anti-mouse IgG.

The supernatant was recovered and passaged twice in Marc-145 cells and identified by reverse transcription PCR. Viral RNA from rHuN4-F112, rHuN4-F112-m1B, rHuN4-F112-m2B, rHuN4-F112-C5, and rHuN4-F112-C5-m1B-m2B were extracted using a QIAamp Viral RNA Mini Kit (Qiagen, Hilden, Germany). Reverse transcription reactions were performed at 2 °C for 10 min and 42 °C for 1 h using the M-MLV
Reverse Transcription Polymerase System (TaKaRa) and then amplified with B-900-F/B-900-R primers (Table 2), and the PCR products were sent for DNA sequencing. Viral rescue was quantified in Marc-145 cells.

**Growth kinetics of deletion-marker viruses in Marc-145 cells**

To monitor the effect of deletion on viral replication, Marc-145 cells were grown to 80% confluency in a six-well plate and then infected with the third-passage viruses (rHuN4-F112-C5 and rHuN4-F112-C5-m1B-m2B) at a multiplicity of infection of 0.01. The supernatant was harvested at 0, 24, 48, 72, and 96 h post-inoculation. After RNA extraction and reverse transcription, the viral copy numbers at different time points were determined by quantitative reverse transcription PCR as described previously (Wei et al. 2008). All samples were tested thrice and the replication kinetic curve was drawn using GraphPad software (GraphPad, Inc., La Jolla, CA, USA). Measured values were expressed as the mean ± standard deviation. Significance was assessed using Student’s *t*-test at *P* < 0.05.

**Results**

**Indel polymorphic analysis of NSP2**

To systematically analyze the indel polymorphism of NSP2, all 907 NSP2 full-length sequences of PRRSV-2 deposited in GenBank between 1991 and 2020 were aligned. Using VR-2332 as a reference strain, extensive indels were found in NSP2 (Fig. 6). They were divided into five main patterns: classic PRRSV (no indels), NADC30 (deletion at positions 322–432, 483, and 504–522), NADC34 (deletion at position 335–434), HP-PRRSV (deletion at positions 482 and 533–561), and SP (36-amino acid insertion at position of 814). All five main indel patterns were detected in PRRSVs isolated in China. During indel analysis, deletions at position 585–586 were found to be somewhat unique. To investigate whether the deletion could be used to design a DIVA vaccine, the antigenic potential of the m1B peptide, whose start-stop sequence spanned positions 562–627 (Fig. 1) and thus covered residues 585–586, was assessed. The same approach was applied with peptide m2B, whose start-stop sequence spanned positions 749–813 and was universal in PRRSV-2.

**Peptides m1B and m2B react strongly with PRRSV-positive serum**

To inspect the antigenicity of m1B and m2B, the corresponding peptides were chemically synthesized and employed to coat an ELISA plate. Pig serum positive for HP-PRRSV or NADC30-like PRRSV was used as the primary antibody. Using indirect ELISA, both m1B and m2B were shown to react strongly with the two sera (Fig. 2A).

**No cross-reaction with other antisera**

The specificity of 1B2B-ELISA indirect ELISA was further examined using positive sera containing antibodies for other swine viral pathogens. No cross-reactions were found, as all values were below the cutoff point (Fig 2B).
Identification of immunodominant antigen regions in m1B and m2B peptides

To identify the antigenic regions with good immunogenicity in peptides m1B and m2B, 14 truncated and overlapping short peptides of m1B or m2B were coated onto ELISA plates, and their immunogenicity was probed with PRRSV-positive serum. The peptides m1B1 (562–577), m1B4 (586–601), m1B5 (594–609), and m1B7 (611–627) were recognized by the positive sera (Fig. 3A, 3B). The truncated peptide m2B5 (781–796) reacted well with NADC30-like PRRSV-positive serum, whereas other m2B short peptides displayed only a weak reaction (Fig. 3C, 3D).

Sequence conservation analysis of m1B and m2B

Sequence alignment of NSP2 revealed that m1B and m2B were conserved within the lineage but diverged from other lineages (Fig. 4). The C-termini of m1B1, m1B5, and m1B7 were relatively conserved among lineages, as was in part the middle portion of m2B5; however, in general, the epitope sequences varied considerably among lineages.

Rescue of PRRSVs harboring m1B or m2B deletions

Using the infectious HuN4-F112 MLV clone as template, deletions of the m1B or m2B regions were generated, and rescue of the corresponding PRRSV deletion mutants (rHuN4-F112-m1B, rHuN4-F112-m2B, and rHuN4-F112-C5-m1B-m2B) was assessed in Marc-145 cells. A cytopathic effect was observed in Marc-145 cells infected with the deletion mutants, as indicated by the bright intracellular immunofluorescence (Fig. 5A). All rescued viruses were sequenced at passage 3, and the specific deletions in NSP2 were confirmed.

Deletion of m1B or m2B does not reduce viral growth

The titers of rHuN4-F112-C5 and rHuN4-F112-C5-m1B-m2B in Marc-145 cells were 10^{5.485} and 10^{5.536} median tissue culture infectious dose per milliliter, respectively. The viral growth curve was assessed by quantitative reverse transcription PCR. The copy number of the double mutant was not significantly different from the parental virus rHuN4-F112-C5 at each time point (Fig. 5B). This result indicates that simultaneous deletion of m1B or m2B at positions 562–627 or 749–813 in NSP2 did not significantly affect the replication of PRRSV in Marc-145 cells.

Discussion

PRRS has caused large economic losses in the pig industry in the past and remains a challenging problem in pig farming. Several killed vaccines or MLVs have been licensed and are widely used. Although these vaccines have played an important role in preventing and controlling PRRS epidemics, they are also associated with some limitations, such as limited immunoprotection against homologous PRRSVs (Hu and Zhang 2014) and the difficulty in distinguishing between vaccinated and infected animals (Corzo et al. 2010). We previously described two amino acid deletions in NSP2 of PRRSV during
viral passage in Marc-145 cells, which suggested that these or nearby residues can be deleted. In the present study, we evaluated the generation of a marker vaccine strain by deleting strongly antigenic gene regions non-essential for viral replication.

NSP2 presents high antigenicity, and many epitopes have been identified in this protein (Bi et al. 2019; de Lima et al. 2006; Yan et al. 2007), including immunodominant B cell epitopes and some potential T cell epitopes (Oleksiewicz et al. 2001). In this study, we identified two immunodominant antigenic regions, m1B and m2B, which reacted strongly to pig serum positive for HP-PRRSV or NADC30-like PRRSV. The sequences of m1B and m2B are not conserved among lineages because of the high genetic variability of PRRSV. We did not further refine the epitopes, mainly because the reactivity of the refined short peptides was not sufficiently strong, as indicated by their positive to negative ratio of around 2. Therefore, rather than using individual epitopes, we combined the m1B and m2B peptides when coating the ELISA plates.

NSP2 is the largest protein of PRRSV, reaching 1196 residues in VR-2332, the prototype strain of PRRSV-2. Insertions and deletions in NSP2 are frequent (Fig. 6), particularly in the middle portion (Han et al. 2007) such as in NADC30-like PRRSVs, NADC34-like PRRSVs, and HP-PRRSVs. This finding suggests that NSP2 can tolerate native amino acid indels of exogenous genes. In theory, indels can be repaired by recombination of two different PRRSV strains; however, no such event has been detected. We analyzed more than 700 PRRSVs-2 deposited in GenBank between 1991 and 2020. High-frequency hotspots of inter-lineage recombination were found in NSP9 and the GP2-GP3 region. In contrast, in NSP2, the frequency of inter-lineage recombination is very low in PRRSVs from China or the United States (Yu et al. 2020). Therefore, the possibility of restoring these mutants to wild-type PRRSV by genetic recombination is very low.

Numerous studies have confirmed the elevated tolerance of PRRSV NSP2 towards indels. This suggests that NSP2 an ideal site for developing recombinant vaccine markers. Han et al. (2007) identified 324–726 amino acids in the middle region of NSP2 as non-essential for replication and whose deletion did not affect virus rescue. In another study using the HuN4-F112 infectious clone as backbone, 25 amino acids were deleted in the non-essential region of NSP2 (at position 508–532), and the dominant epitope of the Newcastle disease virus nucleoprotein (NP49) was inserted in their place to construct a double-marker attenuated vaccine strain (rHuN4-Δ25+NP49) (Xu et al. 2012). The insertion position, size, and nature of the foreign gene may affect the viability and genetic stability of the obtained virus. Among previous attempts to insert genes encoding the green fluorescent protein (GFP), luciferase, and Flag tag in the non-essential region of NSP2, only GFP insertion resulted in the rescue of PRRSV particles. With passage of the virus, the inserted GFP accrued gene mutations and deletions until it lost its activity (Fang et al. 2006; Kim et al. 2007). Thus, the reason why exogenous genes cannot exist in a stable form in NSP2 requires further exploration.

In the present study, two peptides that are universal in NSP2 of PRRSV-2 were found to be immunodominant. This characteristic can be used to monitor antibodies against PRRSV by peptide-coated ELISA. More importantly, PRRSV mutants with deletion of either of the two peptides multiplied
well in Marc-145 cells, suggesting the potential of the two peptides as molecular markers for developing a DIVA vaccine against PRRSV and the accompanying differential diagnosis methodology.

## Declarations

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### Authors’ contributions

ATQ conceived and designed research. LDY, TXX, YYB and HY conducted experiments. CXH, TJJ and HXY contributed new reagents or analytical tools. WT and WQ analyzed data. LDY wrote the paper; ATQ revised the paper. All authors read and approved the final manuscript.

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### Data availability

All the sequences are available from GenBank.

### Ethics approval and consent to participate

Not applicable.

### Conflict of interests

The authors declare that they have no conflict of interests.

### Ethics approval

This article does not contain any studies with human participants or animals performed by any of the authors.

### Consent for publication

Not applicable.

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Tables

Table 1. Amino acid sequences of short-peptides in this study

| Peptide name | Sequence          | Positions in NSP2 |
|--------------|-------------------|-------------------|
| m1B1         | TTTLTHQDEPLDPAS   | 562-577           |
| m1B2         | PLDLPASSQTEYEAFP  | 570-585           |
| m1B3         | SQTEYEAFPLAPSQNM  | 578-593           |
| m1B4         | LAPSQNMGILEAGGQE  | 586-601           |
| m1B5         | GILEAGQEVEEVLSEI  | 594-609           |
| m1B6         | VEEVLSEISDILNDTN  | 602-617           |
| m1B7         | SDILNDTNPAPVSSSSS | 611-627           |
| m2B1         | IGSVATEDVPRILGKIG| 749-764           |
| m2B2         | PRILGKIGDTDELLDR  | 757-772           |
| m2B3         | DTDELLDRGPSAPSKG  | 765-780           |
| m2B4         | GPSAPSKGEPVCDQPA  | 773-788           |
| m2B5         | EPVCDQPAKDPROMSPR | 781-796           |
| m2B6         | KDPRMSPRESDESMIA  | 789-804           |
| m2B7         | ESDESMIAPPADTGGVG | 797-813           |

Table 2. Primers used in this study
| Primer name  | Sequence                          |
|-------------|-----------------------------------|
| 1. B-FseI-F | GAGTGGGCGGGCCGCCCCAGT           |
| 2. B-Nhel-R | GCCGCTGGCGGCTAGCAG            |
| 3. B-m1B-F  | GTGGCGGCAGCTCAAGTGTAAGATCA       |
| 4. B-m2B-R  | CGGCAAATCAGTAAGAAATGGTGAGGTCACCTC |
| 5. B-m2B-F  | H-1B-2B-F: GAGTGACCTCACATTTCATTCACTGATTGCGGTCT  |
|             | H-1B-2B-R: CAGAGATGACCTCACATTTCATTCACTGATTGCGG  |
|             | B-900-F: CGGCAAATCAGTAAGAAATGGTGAGGTCACCTCTCTG |
|             | B-900-R: AGTGGAGCCGTACTTATG      |
|             | ACCCCCATTTCGACACCG             |

Figures

**Figure 1**
Insertions and deletions in NSP2 of circulating PRRSV-2 or MLV strain. VR-2332, the prototype of PRRSV-2, was used as a representative strain. Strains are listed on the left. The positions labeled in the figure are preferred to the corresponding position in VR-2332. Two regions (named m1B and m2B) at 562–627 and 749–813 are universal in PRRSV-2. On the left is the name of the representative strain of each indel type. NSP2, non-structural protein 2; PRRSV, Porcine reproductive and respiratory syndrome virus.

Figure 2
Antigenicity analysis and cross-reactivity assessment of m1B and m2B (A) Indirect ELISA of m1B or m2B peptides. ELISA plates were coated with synthesized peptides, whereas pig serum positive for HP-PRRSV or NADC30-like PRRSV was used as the primary antibody. (B) ELISA plates were coated with synthesized peptides, whereas positive serum for PRRSV, CSFV, PEDV, PRV, PPV and PCV2 was used as the primary antibody. The black dotted line represents the cutoff value of 1B2B-ELISA. PRRSV, Porcine reproductive and respiratory syndrome virus; HP-PRRSV, highly pathogenic PRRSV; CSFV, classical swine fever virus; PEDV, porcine epidemic diarrhea virus; PRV, pseudorabies virus; PPV, porcine parvovirus; and PCV2, porcine circovirus type 2.
Figure 3

Immunoreactivity of m1B and m2B truncated peptides. (A) Schematic diagram of overlapping m1B1–m1B7 short peptides used for epitope mapping. The green box represents peptides that reacted strongly with PRRSV-positive serum, whereas the gray box represents peptides that reacted only weakly. (B) P/N ratio for m1B1–m1B7 in indirect ELISA. (C) Schematic diagram of overlapping m2B1–m2B7 short peptides. (D) P/N ratio for m2B1–m2B7 in indirect ELISA. P/N, Positive to negative.
Figure 4

Sequence analysis of m1B and m2B. Representative strains of current circulating PRRSVs or MLVs are shown. The positions of indels in m1B and m2B are labeled below the sequences, whereas overlapping short peptides (m1B1–m1B7 and m2B1–m2B7) are labeled above the sequences. A purple line indicates strong reactivity between the short peptide and PRRSV-positive serum, whereas a gray line denotes no reactivity. PRRSV, Porcine reproductive and respiratory syndrome virus; MLV, modified live vaccines.
Figure 5

Characterization of PRRSV m1B and m2B deletion mutants. (A) CPE and bright fluorescence of deletion marker strains and parental PRRSV infected MARC-145 cells. a, rHuN4-F112-m1B; b, rHuN4-F112-m2B; c, rHuN4-F112-C5-m1B-m2B; d, rHuN4-F112; e, rHuN4-F112-C5; f, negative control. Bar: 200 µm. (B) Growth curves of the parental virus and deletion mutants in Marc-145 cells. Values represent the average of three independent experiments. PRRSV, Porcine reproductive and respiratory syndrome virus; CPE, Cytopathic effect.
Figure 6

Systematic indel patterns of NSP2 based on 907 PRRSV-2. Indel patterns of NSP2 are divided into five main categories. The arrows and numbers indicate the indel locations in NSP2 using the position in the VR-2332 strain as reference. Values on the right represent the numbers of strains. NSP2, non-structural protein 2; PRRSV, Porcine reproductive and respiratory syndrome virus.