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Peptides with 16R in S2 protein showed broad reactions with sera against different types of infectious bronchitis viruses

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**ABSTRACT**

Vaccination plays a vital role in controlling diseases caused by chicken infectious bronchitis virus (IBV). The continuously variant antigenicity of IBV limits the application of current vaccine strategies and serological diagnostic systems. S2 protein is an invariant that harbors broad neutralizing epitopes. However, little is known about the key amino acids that contribute to the broad-spectrum S2 epitopes. In this study, we aimed to elucidate the specific amino acids contributing to S2 epitopes. Site mutagenesis and peptide-based enzyme-linked immunosorbent assays (ELISAs) showed that 16R in S2 protein was a key amino acid mediating the antigenicity of S2 protein. S2-derived peptides with 16R, but not those with 16 K, could react with sera against different types of IBVs. Notably, a commercial ELISA kit for detection of antibodies against IBV did not react with sera against all types of IBVs. Taken together, these data demonstrated that S2-derived peptides with 16R could be used as novel marker-based antigens for developing both broad-spectrum vaccines and serological diagnostic kits to control IBV.

1. Introduction

Infectious bronchitis (IB) is a highly contagious disease caused by IB virus (IBV). IBV infection generally causes serious respiratory and renal diseases in meat chickens and decreased egg laying in laying hens (Cavanagh, 2007), resulting in huge economic losses in the poultry industry (Bande et al., 2017). Although IBV vaccines are widely used, wild-type IBV or IBV escape mutants are frequently isolated from vaccinated chicken flocks because of the circulation of multiple serotypes in flocks and continuous mutation of the protective epitopes of IBVs. Thus, current vaccine strategies need to be re-evaluated, and more efficient, broadly protective IBV vaccines or diagnostic approaches are urgently needed.

The IBV genome encodes four major structural proteins, i.e., spike (S), small envelope (E), membrane (M), and nucleocapsid (N), as well as 15 nonstructural proteins and some accessory proteins (Lin and Chen, 2017). Among these proteins, S glycoprotein is the major protective antigen carrying neutralizing epitopes and can induce efficient immune responses against IBV (Ignjatovic and Galli, 1993, 1995; Kant et al., 1992). Notably, S protein can be cleaved into the amino-terminal S1 and carboxy-terminal S2 domains by a furine-like protease from the host (Cavanagh, 2007; Eldemery et al., 2017; Kusters et al., 1989). S1 is responsible for IBV attachment to host cells, whereas S2 contributes to viral fusion activity of IBV (Wickramasinghe et al., 2014). Both S1 and S2 play vital roles in mediating IBV infection in host cells. S1 is a major protective antigen for IBV, but is highly variable among different IBV stains (Ignjatovic and Galli, 1994). In contrast to S1, S2 is highly conserved and carries broad antigenic epitopes (Toro et al., 2014). Therefore, S2 could be a marker for developing more efficient vaccines and broad diagnostic approaches.

Previously, S2 glycoprotein from IBV was not thought to induce virus-neutralizing antibodies or chicken tracheal protection (Cavanagh et al., 1986). (Ignjatovic and Galli, 1995) found that S2 glycoprotein was the most immunogenic structural protein following vaccination with inactivated virus. S2 protein expressed from recombinant virus confers broad protection against challenge (Toro et al., 2014). Therefore, S2 protein has important roles in inducing protective immunity (Eldemery et al., 2017). Moreover, (Kusters et al., 1989) first identified a conservative immunodominant region near the N-terminus of S2 protein. Recently, several regions (e.g., SNCPLVSYGKFCIKPDGSI27, 2019 Elsevier B.V. All rights reserved.
designated SE8-27) in S2 have been identified as broad or neutralizing epitopes for IBV (Andoh et al., 2018; Ignjatovic and Sapats, 2005). However, little is known regarding the key amino acids that contribute to broad-spectrum S2 epitopes.

In this study, we analyzed 330 IBV isolates from GenBank database and evaluated effects of mutations in the SE8-27 epitope in S2 on antigenicity.

2. Materials and methods

2.1. Virus and serum samples

M41 strain (GenBank accession number: DQ834384; mass type IBV) and 4/91 strain (GenBank accession number: KF377577; 4/91 type IBV) of IBV was obtained from Sinopharm Yangzhou Vacc Biological Engineering Co., Ltd. (Yangzhou, China). CK/CH/2010/JT1 (GenBank accession number: KU361187; new cluster type IBV), CK/CH/2014/QL1403 (GenBank accession number: KU361198; TC07-2 type IBV), and CK/CH/2014/FJ14 strains of IBV (GenBank accession number: MN262521; QX-like type IBV) were isolated and identified by our laboratory. SPF white leghorn chicken eggs were purchased from Beijing Merial Vital Laboratory Animal Technology Co., Ltd. (Beijing, China) and hatched out in our laboratory. All birds were kept in isolators throughout the experiments. To prepare serum against M41, 4/91, CK/CH/2010/JT1, CK/CH/2014/QL1403, and CK/CH/2014/FJ14 strains, 14-day-old SPF chickens were inoculated intraocular-nasally with 10^3–10^5 EID_{50} of different strains. Thirty days after immunization, serum from immunized chickens was collected and stored at −20°C. All experiments complied with institutional animal care guidelines and were approved by the University of Yangzhou Animal Care Committee.

2.2. Preparation of the peptide library

Amino acid sequences of the S2 protein of IBV strains, including H120, D207, and CK/CH/2010/JT1 strains, were obtained from GenBank (accession numbers: MK937831, X58003, and KU361187, respectively). Six peptides (Pep1–Pep6) with good conservation and antigenicity were selected by MegAlign and Protean software, version 7.1.0 (DNAstar). Pep1 is a conservative epitope for all IBV strains (Ignjatovic et al., 2005). Pep2–Pep5 were designed according to conserved peptides in the S2 gene of IBV. Peptides containing between 16 and 20 amino acid residues were synthesized by Synpeptide Co., Ltd. (Shanghai, China). The peptide sequences are listed in Table 1.

2.3. Enzyme-linked immunosorbent assay (ELISA)

For ELISA, 96-well plates were coated with 0.1 μg/well synthetic peptide in 0.1 M carbonate buffer (pH 9.6) at 4°C overnight (Khairy et al., 2017). After washing with phosphate-buffered saline (PBS) with Tween 20 (PBST), the plates were blocked with 300 μl of 8% rabbit serum (Lanzhou Minhai Biological Engineering Co., Ltd., China) for 3 h at 37°C. Following three washes with PBST, 100 μl serum (1:200 in PBST) was added to the wells, and the plates were incubated at 37°C for 1 h. The plates were then washed five times and further incubated with 100 μl horseradish peroxidase (HRP)-conjugated rabbit anti-chicken IgG (Jackson ImmunoResearch Laboratories, Inc., USA) at 1:30000 dilution in blocking buffer for 1 h at 37°C. After washing five times, and the colorimetric reaction was developed by incubating the plates with 100 μl TMB substrates for 15 min at 37°C. Color development was stopped with 100 μl of 1% sodium dodecyl sulfate. The absorbance at OD_{450} was obtained using an ELISA reader (BioTek, VT, USA). There were five serum samples for each genotype IBV strain. Each experiment was repeated twice.

A commercial IBV antibody detection kit was purchased from IDEXX (USA). Briefly, all reagents were allowed to warm to 18–26°C before use. Negative and positive controls were dispensed into duplicate wells. Samples diluted 500-fold with sample diluent were added to appropriate wells and incubated for 30 min. The plates were washed with approximately 350 μl distilled water five times. Next, 100 μl HRP-conjugated secondary antibodies was added to each well, and plates were incubated for 30 min. After washing five times, 100 μl TMB substrate was dispensed into each well, and plates were incubated for 15 min. Color development was stopped with 100 μl stop solution. The absorbance at OD_{450} was then determined. Positive results were defined as S/P values over 0.2.

2.4. Sequence analysis of IBV S2 protein

All available 330 complete genome IBV strains were downloaded from the NCBI database. The sequences were analyzed using MegAlign and Protean software, version 7.1.0 (DNAstar).

2.5. Statistical analysis

All the data were graphed, and statistical analyses were performed using Prism 5 software (GraphPad, La Jolla, CA). One-way ANOVA with repeated measures was used to evaluate the reactivity of peptide antigens with sera against IBV. The differences were considered statistically significant at p values of < 0.01.

3. Results

3.1. Variations in the broad-spectrum region (amino acids 8–27) in the S2 protein of IBV

The region containing 8NCPYVSYGKFCIKPDGSIST27 in S2 was downloaded from GenBank. After analysis of complete genome sequences of 330 IBV isolates, we found that almost all IBV strains contained a K at amino acid position 16 (16K) in S2 before the 1990s. Subsequently, strains with an R at amino acid position 16 (16R) emerged and became the dominant pandemic IBV strains. Interestingly, 16K IBV strains were mainly associated with respiratory tract diseases, and the 16R IBV strains were mainly associated with renal pathogenicity. Based the region with 8NCPYVSYGKFCIKPDGSIST27 in S2, IBV strains could be divided into two groups: Group I containing 16K, and Group II containing 16R (Table 2). Two strains, i.e., B1648 (accession number: KR231009) and gamaCoV/ck/China/I0636/16 (accession number: MH924835), contained T and Q as the 16th amino acids (group I-like), respectively.
To elucidate which single amino acid was essential for the antigen reaction of CK/CH/2010/JT1 strain in this region, Pep6 was synthesized according to the same region in S2 in the CK/CH/2010/JT1 strain. Pep6, which contained the epitope of Group II, reacted with all sera against the five different types of IBVs, whereas Pep1, which contained the epitope of Group I, only reacted with four of five different types of IBVs (Fig. 1A, 1B). Pep1 could not efficiently react with sera against IBV CK/CH/2010/JT1 (new cluster type IBV). Pep4 and Pep5 could react with sera against M41 type and 4/91 type, whereas Pep2 and Pep3 only reacted with sera against M41 type and 4/91 type, respectively. Notably, a commercial ELISA kit for detection of antibodies against IBV was also used to detect sera against different types of IBVs. The commercial kit could only react with sera against M41 and 4/91 strains, but not with sera against CK/CH/2014/QL1403, CK/CH/2014/FJ14, and CK/CH/2010/JT1 strains (Fig. 1C).

3.3. 16R was identified as a key amino acid for S2 broad antigenicity

To identify the critical amino acids contributing to the broad-spectrum antigenicity of Pep6, the last four amino acids were deleted in other peptides because they showed high variation in different genotype or the same genotype of IBV. We used Pep1-1 as a prototype and Pep1(epitope SE8-27) and Pep6 (amino acids 8–27 from S2 of Group I and Group II, respectively) and Pep2–5 (derived from conserved peptides of IBV S2 protein; Table 1) were synthesized and used as peptide antigens for testing crossreaction with sera from different genotypes of IBVs, including mass, QX-like, TC07-2, 4/91, and new cluster types. We analyzed the reactivity between the peptides and different genotypes of IBVs. Only Pep1 showed good reactions with most sera. However, Pep1 did not react with the sera from CK/CH/2010/JT1 strains. After comparing and analyzing amino acid sequences from different genotype IBV strains in this region, we found that there were some differences in amino acids among different genotype strains.

To further evaluate and identify the broad antigenic epitopes in S2, 16R was identified as a key amino acid for S2 broad antigenicity. The black/red dotted lines represent critical values. There were five serum samples for each genotype. Each experiment was repeated twice. The results show the mean values. (A) Reactivity of Pep1–Pep6 with sera against different IBV strains. (B) Sequences of amino acids 8–27 in S2 protein from different IBV strains and reactivities of Pep1–Pep6 in sera against different IBV strains. “+” represents peptides that reacted with sera harboring IBV; “−” represents peptides that did not react with sera harboring IBV. (C) Reactivity of a commercial ELISA kit with sera against different IBV strains. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

3.2. Pep6, not Pep1, showed broad reaction with sera from different types of IBVs

To further evaluate and identify the broad antigenic epitopes in S2, the epitope of Group I (136/328) and Pep6, not Pep1, showed broad reaction with sera from different types of IBVs. Only Pep1 showed good reactions with most sera. However, Pep1 did not react with the sera from CK/CH/2010/JT1 strains. After comparing and analyzing amino acid sequences from different genotype IBV strains in this region, we found that there were some differences in amino acids among different genotype strains.

Table 2

Sequence analysis of amino acids 8–27 in S2 protein from 314 IBV strains.

| Genotypes | Sequences of amino acids 8–27 in S2 protein |
|-----------|--------------------------------------------|
| Group I   |                                            |
| GI-1, GI-2, GI-3, GI-4, GI-5, GI-6, GI-8, GI-9, GI-11, GI-12, GI-13, GI-14, GI-16, GI-21, GI-22, GI-26, GI-1, GV-1 | NCPYYSYGRCKIPDGS Requests: |
| Group II  |                                            |
| GI-7, GI-19, GI-22 | SCSYYSYGRCKIPDGS Requests: |

Notes: Red highlighting indicates differences in amino acid 16, which were used for grouping.

Fig. 1. Reactivity of Pep1–Pep6 with sera against different IBV strains. The black/red dotted lines represent critical values. There were five serum samples for each genotype. Each experiment was repeated twice. The results show the mean values. (A) Reactivity of Pep1–Pep6 with sera against different IBV strains. (B) Sequences of amino acids 8–27 in S2 protein from different IBV strains and reactivities of Pep1–Pep6 in sera against different IBV strains. “+” represents peptides that reacted with sera harboring IBV; “−” represents peptides that did not react with sera harboring IBV. (C) Reactivity of a commercial ELISA kit with sera against different IBV strains. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
mutated N8S, K16R, and K20Q, yielding Pep1-1-1, Pep1-1-2, and Pep1-1-3, respectively. The results demonstrated that 16R in Pep6-1, not 8S or 20Q, contributed to the broad-spectrum antigenicity of this epitope (Fig. 2A). Pep1-1-2 with 16R (similar to Pep6) could react with all sera against different types of IBVs (Fig. 2C). Pep1-1-4, in which amino acid 16 (R) was mutated to K, similar to Pep6-1, did not react with positive sera against CK/CH/2010/JT1 strain. Taken together, these results showed that 16R was the only determinant of the broad-spectrum antigenicity of Pep6.

3.4. Distribution of Group II strains carrying 16R in the epitope

Next, we downloaded and analyzed all complete genome sequences of IBVs reported in the NCBI database (https://www.ncbi.nlm.nih.gov/). The information on the geographical origin and time of isolation of these sequences were also obtained from the NCBI database. Sequence analysis further revealed that IBV strains carrying 16R in the epitope of Group II were first isolated in the 1990s, and the proportion then increased thereafter (Fig. 3A). IBV strain KM91, which was isolated in 1991, was the first IBV with such an epitope. The incidence of 16R-containing IBV strains increased over time. Variations in key amino acids in the conserved S2 protein suggested that the IBVs had undergone considerable variation and evolution, necessitating new vaccine strategies and diagnostics to prevent and control IBV. Currently, the epitope of Group II is widely present in IBV isolates from Asia, America, Europe, and Africa (Fig. 3A, 3B), consistent with the prevalence of the lineage GI-19, the so-called QX-type IBV, in China and worldwide (Zhao et al., 2017). Our results demonstrated that the epitope of the QX type IBV dominant in chicken flocks in China all belonged to this group of IBV strains containing 16R.
4. Discussion

IBV is a typical coronavirus that is constantly recombining and mutating its antigenicity, leading to the presence of multiple serotypes and genotypes of IBV strains and causing major challenges in disease prevention and control (Cavanagh, 2007). As an important antigenic protein, S1 has been used in vaccine development (Hodgson et al., 2004; Wickramasinghe et al., 2014; Zou et al., 2015). Notably, S2 protein also can induce protective immune responses (Eldemery et al., 2017; Ignjatovic and Sapats, 2005). In contrast to S1, S2 protein is thought to be invariant and contain neutralizing epitopes (Wickramasinghe et al., 2014). Of these epitopes in S2, the epitope containing amino acids 8–27 has been identified as a broad antigenic epitope for IBVs. Our data demonstrated that this epitope in S2 protein from different types of IBVs was also not highly conserved in different IBV serotypes and could be divided into two groups, designated Group I (8NCYPVSYKFC1KPDGSI2T7-like) and Group II (SCPYPVSYGRFCIQ-PDGSIKQ27-like). The two strains, B648 (accession number: KR231009) and gamaCoV/ck/China/10636/16 (accession number: MH924835), were classified as Group I because their 16th amino acids were T and Q, respectively.

To identify the key amino acid in Pep6, the last four amino acid of Pep1 and Pep6 were removed, as previously described (Kusters et al., 1989). Our results also showed that the last amino acid had minor effects on antigenicity. Further mutational analysis confirmed that mutation of the 16th amino acid (X16R) was the key for the broad-spectrum antigenicity of this epitope.

Serological assays further showed that the S2-derived epitope in Group II, but not that in Group I, could react with all sera against different types of IBVs. Moreover, 16R was identified as a key amino acid for the broad antigenicity of Group II. Valastro et al. (2016) performed a phylogenetic analysis based on the S1 gene of 1286 strains to derive a new and coherent classification scheme for IBVs. They divided the IBV strains into six genotypes, labeled GI, GII, GIII, GIV, GV, and GVI. GI contained 27 lineages, whereas GII, GIII, GIV, GV, and GVI contained only one lineage each. Our grouping method was based on S2 protein of IBV. We further compared two different grouping methods and found that GI-7, GI-19, and GI-22 belonged to Group II, whereas the other strains belonged to Group I. Notably, widespread XQ genotype, new cluster genotype, TW-1 type, LDT3 type, and BJ type IBVs all carried 16R in the S2 protein. This suggested that peptides with 16R derived from S2 protein could be used as novel marker-based antigens for developing both broad-spectrum vaccines and serological diagnostic kits to control IBV. Whether the 16R in the IBV S2 protein playing a role in tissue tropism and induced immune response remains to be investigated in further.

Declaration of Competing Interest

The authors declare that they have no competing interests.

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