Introduction

Newcastle disease virus (NDV) is a member of the avian paramyxovirus (APMV) group belonging to the *Avulavirus* genus in the family *Paramyxoviridae* [26]. There are currently nine known antigenic serotypes of APMV, designated APMV-1 through -9. NDV is the sole member of APMV-1 and is an enveloped virus with a negative-sense, single stranded RNA genome which codes for six structural and two non-structural proteins [7,14]. The structural proteins include nucleocapsid (N), phospho-(P), and matrix (M) proteins, fusion (F) and hemagglutinin-neuraminidase (HN) glycoproteins, and a large polymerase (L). Of these, the HN and F proteins are membrane anchored viral glycoproteins that enable cell binding and infection. The HN glycoprotein recognizes sialic acid containing receptors on cell surfaces and acts as a neuraminidase, removing sialic acid from progeny virus particles. The F protein is presumed to mediate membrane fusion either at the cell surface [8] or through receptor-mediated endocytosis [5].

Newcastle disease (ND) caused by virulent ND virus is one of the most devastating infectious diseases of poultry and induces up to 100% mortality as well as a decrease in egg production and poor egg quality [4]. NDV strains are classified into lentogenic, mesogenic, and velogenic categories, a nomenclature which reflects increasing levels of virulence for chickens based on biological parameters such as mean intracerebral pathogenicity index.

The molecular determinant for NDV virulence is known to be related to the cleavability of precursor F (F0) to active F1 and F2 polypeptides by cellular proteases [13,18,23]. Virulent NDVs have a multibasic amino acid sequence (i.e. 112RRQKR117) at the F0 cleavage site (virulence motif), whereas low virulent NDVs have a monobasic amino acid sequence (i.e. 112GRQGRL117) in the same region (avirulence motif) [2,6,21]. Most of the recent virulent NDV strains bear the virulence motif 112RRQKR117 at the cleavage site of their F0 protein [1,15,17,21].

Polyclonal anti-peptide sera to peptides whose sequences mimic the F2-polypeptide C-termini (i.e. 112RRQ115) have been used to detect specifically virulent NDV isolates [12,16,25]. Despite the usefulness of its diagnostic applications, the antigenic and immunogenic properties of the motif still remain unclear. In this study, we map a B-cell epitope of the virulence motif 112RRQKR117 and investigate whether the antibody response to the B-cell epitope on the virulence motif is induced in virulent NDV-infected chickens.
Materials and Methods

Synthetic peptides
The synthetic peptide RRQRKF (CAARRQKFRI) containing the virulence motif RRQRKF was synthesized by a commercial company (Peptron, Korea) using Fmoc (9-fluorenylmethoxycarbonyl) solid-phase synthesis chemistry on Rink amide resin (Novabiochem, USA). The peptide was subjected to C-terminal amidation to stabilize its structure. Six peptides were prepared by an alanine substitution of the 112RRQRKF 117 motif with at each position (e.g. R112A represents an arginine to alanine substitution at residue 112; Fig. 1). Four synthetic peptides containing other virulence motif (KRQKRF, RRQRKF and RRRKRF) or avirulence motif (GRQGRL) sequences were also synthesized in the same manner. Each peptide was then conjugated to bovine serum albumin (BSA) for enzyme-linked immunosorbent assay (ELISA).

Anti-RRQKRF antibodies
Anti-RRQKRF monoclonal antibody (mAb) 4G2 was used in this study. The hybridoma cell line producing mAb 4G2 (IgG1, kappa) was generated from mice immunized with the keyhole limpet haemocyanin (KLH)-conjugated synthetic peptide CGGRRQRKRFI in our laboratory. Ascitic fluids were produced following inoculation of 107 hybridoma cells into the peritoneal cavities of pristine-primed Balb/c mice as described elsewhere.

Anti-RRQKRF sera were prepared from chickens (n = 2) by two subcutaneous inoculations with the KLH-CAARRQKFRI peptide conjugate (100 μg per bird) at three week intervals. Sera were taken from immunized birds 3 weeks after the last inoculation. Sera were diluted in pre-immune chicken sera and then titrated by ELISA. Diluted serum having 25% of the highest optical density (OD) value in ELISA was used as positive serum control. Pre-immune chicken sera served as negative serum control.

Virus
NDV strain Kr-005/00 was used in this study. This strain is a highly virulent NDV strain having the virulence motif 112RRQRKF 117 at the cleavage site of the F0 protein [15]. NDV was propagated by using SPF chicken eggs, and stored at −70°C until use.

Animal experiment
Six week-old specific pathogen-free (SPF) chickens from White Leghorn parents (Lohmann, Germany) were used in this study. The SPF birds were maintained in air-filtered bio-security isolation units (ThreeShine, Korea) with feed and water ad libitum for the duration of the study. All animals were kept at animal facilities and we followed the ethical guidelines of the Animal Welfare Committee recommended by the National Veterinary Research and Quarantine Service during the experiment. A total of 24 six-week old SPF chickens in this experiment were divided into two groups for the experiment. The infection group was challenged with Kr-005/00 (104.5 EID50/100 μL) (n = 12, 100 μL per bird) via the eye drop and intranasal (ED/IN) routes. Control birds (n = 12) were sham challenged with the same volume (100 μL) of PBS via the ED/IN route. All birds were monitored daily for overt clinical signs (depression, respiratory signs, diarrhea etc.) and mortality. Serum and cloacal samples were taken twice on days 0 and 7 post-inoculation.

Virus re-isolation
The swab samples collected for virus isolation were placed in 3 mL transport medium (Micro Test M4RT; Remel, USA) and stored at −70°C until use for virus isolation. For this, the samples were first clarified by centrifugation at 3,000 g for 10 min and then 100 μL of the supernatant was inoculated into allantoic cavity of 9 to 11 day-old SPF embryonated chicken eggs.

Hemagglutination inhibition (HI) test
Serum titers to NDV were determined by HI tests in V-bottom microtiter plates as described elsewhere [3]. Thus, two-fold dilutions of test sera in PBS, pH 7.4, were mixed with the same volume containing 4 HA units of NDV HA antigen (Daesung Microbiological Laboratory, Korea). The mixture (50 μL per well) was then incubated at room temperature for 30 min. An equal volume of 0.5% chicken RBC in PBS was added. The HI endpoint was determined.
as the last dilution that showed complete inhibition of HA activity. All tests were repeated twice. The HI titers are expressed as reciprocal log2 in this study.

**Peptide based ELISA (pELISA)**

The BSA (diluted in 0.01M PBS, pH7.4) was coated on an ELISA plate (Maxisorp; Nunc, France) at 37°C for 2 h. Each serum or mAb sample (50 μL per well) was incubated in two test wells in the ELISA for 1 h. Here, sera and mAb 4G2 were used at dilutions of 1:100 and 1:2,500, respectively, in blocking buffer (0.2% Tween 20 and 2% BSA in 0.01 M PBS). The antigen-antibody reaction was visualized by use of horseradish peroxidase-conjugated 5D2 antibody (BioNote, Korea) or horseradish peroxidase-conjugated anti-mouse IgG (H + L) antibody (Kirkegaard & Perry Laboratories, USA) (100 μL/well, diluted 1:1,500 in blocking buffer) and then TMB (3, 3´, 5, 5´-tetramethylbenzidine) solution (Sigma, USA). ELISA OD was measured at a wavelength of 450 nm. The net OD of each serum was calculated by subtracting the OD value of the control negative well from the mean OD value of the peptide-coated wells. The OD of each well was converted to a percent positive (PP) value of mAb bound to serum antibodies using the following formula: PP = (net OD of test serum/net OD of positive control serum) × 100. Sera having a PP value of 20% or more were considered positive.

**Field samples**

The sera (n = 93) taken from surviving chickens on five farms with ND outbreak were used in this study (Table 1). At the time of sampling, virulent NDV (bearing the RRQKRF motif) was isolated from tissue samples (trachea, cecal tonsil and kidney) from the dead birds (data not shown). For comparison, normal sera (n = 40) taken from apparently healthy chickens on two farms with no ND outbreak were also used. All chickens had received at least two vaccinations with live NDV vaccines (1 and 10 to 14 days) before bleeding. All sera were tested by pELISA using synthetic peptide bearing RRQKRF motif. The results obtained by pELISA were compared with the HI test.

**Statistical analysis**

The pELISA and HI data from serum samples are presented as mean ± SD. Differences in the antibody titers between groups were analyzed by using the Mann-Whitney U test. Significance was accepted at p < 0.05.

**Results**

**Epitope mapping of virulence motif of NDV F protein using monoclonal antibody 4G2**

Virulence motif specific-monoclonal antibody 4G2 was used for mapping B-cell epitope on virulence motif112RRQKRF117. The peptide antigens (RRQKRF, R112A, R113A, Q114A, K115A, R116A and F117A) were tested at a concentration of 4 μg/mL by indirect ELISA using mAb 4G2. As expected, the 4G2 antibody reacted strongly with peptide RRQKRF. It also reacted strongly (ELISA absorbance of >1.5) with peptides R112A, R113A and Q114A, and weakly (ELISA absorbance of 0.68) with peptide K115A. It did not react detectably (ELISA absorbance of <0.2) with peptides R116A, F117A and BSA alone (Fig. 1). This indicates that the 4G2 antibody recognizes the amino acid sequence of the 115KRF117 sequence in the virulence motif of the F protein.

**Reactivity of RRQKRF motif to sera from NDV infected chickens**

The immune response to the virulence motif was then...
examined by pELISA using the BSA-conjugated peptide RRQKRF and sera obtained from virulent NDV infected SPF chickens. SPF chickens were challenged with virulent NDV Kr005 with RRQKRF motif. Prior to the NDV challenge, none of the birds exhibited clinical signs of overt ND and the virus could not be isolated from oral/cloacal swabs. On the day of inoculation, all of the birds had HI antibody titers of < 2 and ELISA absorbance of < 0.2. All the virulent NDV-infected birds displayed severe depression between 2 and 5 days postinoculation (dpi). Sixty seven percent (8/12) of affected birds died showing severe depression between 3 and 5 dpi. The remaining four birds (chicken Nos. 5-1, 5-2, 5-5 and 5-12) survived up to 7 dpi. They were humanely sacrificed after sampling (sera and cloacal swabs) 7 dpi. Virulent NDV was re-isolated 7 dpi from all of the affected birds when cloacal swab samples were inoculated into SPF embryonated chicken eggs. No virus was isolated from cloacal swabs taken 7 dpi from mock-infected control birds.

Sera taken 0 dpi and 7 dpi from infected and control chickens were employed to detect the peptide-bound antibodies. Anti-NDV antibodies were detected in all of the four birds surviving to 7 dpi and their average HI titers ranged from 32 to 256. The 7 dpi sera of the four chickens (5-1, 5-2, 5-5 and 5-12) had significantly increased ELISA PP values (PP value of ranging from 84 to 314) compared to the 0 dpi sera (PP value of ranging from 2 to 14) (p < 0.05) (Fig. 2). All control chickens (n = 12) had HI antibody titers of < 2 and ELISA PP values of < 20 during the experiment.

Reactivity of other virulence motif to sera from NDV infected chickens

We next used pELISA to examine how sera obtained 7 dpi from virulent NDV infected chickens reacted to four synthetic peptides containing three virulence motif (KRQKRF, RRQRRF, and RRRKRF) or avirulence motif (GRQGRL) sequences. Four 7 dpi sera (chicken Nos. 5-1, 5-2, 5-5 and 5-12) were tested. A peptide containing avirulence motif was also included. All of the pre-infection (0 dpi) sera (HI titers of < 2) had a mean absorbance of < 0.2 for peptides KRQKRF, RRQRRF, RRRKRF and GRQGRL. However, the 7 dpi sera for the three peptides KRQKRF and RRQRRF, RRRKRF showed significantly increased ELISA absorbance compared to the 0 dpi sera (p < 0.05); 1.64 ± 0.05 vs. 0.10 ± 0.05 for peptide RRQRRF, 1.11 ± 0.08 vs. 0.10 ± 0.05 for peptide RRRKRF and 2.01 ± 0.15 vs. 0.10 ± 0.19 for peptide RRQKRF (Fig. 3). However, peptide GRQGRL had ELISA absorbance of 0.2 with the 7 dpi sera as well as the 0 dpi.

Reactivity to sera taken from surviving chickens on NDV outbreak farms

We tested the sera taken from surviving birds (n = 93) on five ND outbreak farms and apparently healthy birds (n = 40) on vaccinated farms without ND outbreak by pELISA (Table 1). Synthetic peptide bearing RRQKRF virulence motif was used as antigen for pELISA. The pELISA positive signals (PP value > 20%) were detected in all of five ND outbreak farms. The ELISA-positive rates of farms A, B, C, D and E were 90% (18/20), 91% (20/22), 65% (13/20), 60% (9/15) and 88% (14/16), respectively. When the sera taken from farms A and B were subjected to the HI test, all except

![Fig. 2. Reactivity of synthetic peptide RRQKRF to sera obtained from four chickens 0 and 7 days post-infection (dpi) with a virulent Newcastle disease virus (NDV) strain. Four (5-1, 5-2, 5-5 and 5-12) of 12 infected birds with virulent NDV survived 7 dpi. Surviving birds were tested 0 and 7 dpi. ELISA results of each serum tested was compared to HI titer (log2). Serum having PP value ≥ 20% in the pELISA was considered positive.](image-url)

![Fig. 3. Reactivity of anti-NDV chicken sera to synthetic peptides mimicking four virulence motifs (KRQKRF, RRQRRF and RRRKRF) and one avirulence motif (GRQGRL), as determined by indirect ELISA. These sera were obtained from four chickens 0 and 7 dpi after they were challenged with a virulent NDV strain.](image-url)
two of the ELISA positive samples of farm B also had HI antibody (HI titer of ≥ 2) to NDV. ELISA negative sera (2 for farm A and 2 for farm B) were also HI antibody negative. In commercial farms F and G without ND outbreak, none but one of the sera reacted with synthetic peptide in the pELISA even though 85% and 55% of the sera tested positive for HI test.

### Discussion

Our epitope mapping of virulence motif 112RRQKRF117 revealed that the mAb 4G2 specific for the virulence motif was directed to 115KRF117. Synthetic peptides RRQRRF, revealed that the mAb 4G2 specific for the virulence motif the continuous sequence 115K/R-R-F117 is not likely to form carboxypeptidase from the F2 C-termini [10,11,19]. Thus C-termini of the F2 subunit to locate closely with F117 at the N-termini of the F1 subunit. It is therefore more feasible that the paramyxovirus are linked to each other by a disulfide bond [9]. This allows the basic amino acids (R/K112-R113) at the C-termini of the F2 subunit to locate closely with R117 at the N-termini of the F1 subunit. It is therefore more feasible that such cross-linkage between F1 and F2 subunits may result in a discontinuous conformational epitope R/K112-R113-X-X-X-R117.

Diagnostically, polyclonal anti-peptide sera to peptides whose sequences mimic the F2 C-termini (i.e. 112RRQ114) have been used to capture virulent NDV isolates even though a defined amino acid sequence of the epitope has yet to be determined [12,16,25]. However, a reported limitation of these anti-peptide antisera was that they occasionally did not react with some NDV isolates even with 112RRQ114 motif sequences [16,25]. The reason for this is unclear, although anomalous cleavage/activation process was cited as a reason for their lack of reactivity [16,25]. As mentioned above, the KRF section of the virulence motif can evoke antibody response, suggesting that anti-KRF antibody might react with virulent NDV strains. Thus, use of anti-peptide anti-sera to a peptide mimicking the entire virulence motif RRQKRF might broaden the spectrum of virulent NDV isolates to be captured in immunoassays than previously used anti-peptide antisera to RRQ (the F2 C-termini) alone.

Currently, commercially available vaccines (both live and killed) that are produced using lentogenic NDV strains are widely used for controlling and preventing ND. Because of this, anti-sera to the virulence motif have had the potential to rapidly differentiate virulent NDV isolates and avirulent NDV strains (i.e. lentogenic vaccine strains) using antigen detection assays [12,16,25]. In this study, the 7 dpi sera obtained from infected chickens with virulent NDV (bearing RRQKRF motif) cross reacted with synthetic peptides containing other virulence motifs but not avirulence motifs. This also implies that the virulence motif could be applied for pathotype-specific serologic immunoassay if antibodies bound to the motif are specifically and sensitively detected in NDV infected chickens. If so, the virulence motif would be a target for a serological test that allows infected birds to be distinguished from vaccinated birds.

For this purpose, we performed pELISA based on the synthetic peptide bearing the virulence motif (RRQKRF) to determine whether pELISA could differentiate infected birds from vaccinated chickens under field conditions. As shown in Table 1, most of the sera had HI antibody regardless of virulent NDV infection since all birds had received vaccination two times or more. However, there was a significant difference (p < 0.05) in pELISA reactivity between ND outbreak farms (60–91%) and farms without ND outbreak (0–5%). This suggests that most birds with ELISA positive results on ND outbreak farms produce antibodies to the RRQKRF motif following virulent NDV infection, as demonstrated in SPF chickens after challenge with virulent NDV. In contrast, the ELISA-positive signal in a bird on farm F is presumably non-specific considering that the bird was still apparently healthy at that time of sampling.

In conclusion, we were the first to examine whether pELISA can discriminate virulent NDV-infected birds from birds that have been vaccinated with whole virus vaccines. Our findings indicate that serological testing using a virulence motif has the potential to differentiate infected from vaccinated birds. This suggests that tests like the pELISA may be suitable as a differentiation between infected and vaccinated animals (DIVA) serological tests.
that can be used to monitor herds vaccinated with whole virus vaccines, perhaps in combination with antigen and/or virus isolation methods. It should be noted that while we did not test the ability of the pELISA to detect virulent NDV strains with different virulence motifs such as KRQKRF, we found that the RRQKRF-specific antibodies could cross-react quite well with these alternate virulence motifs. Thus, it may be that antibodies raised against the alternate virulence motifs exhibit similar cross-reactivity. Further studies on development of a virulence motif-based immunoassay for DIVA serological test are in progress.

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