Epidemiological Analysis and Genetic Characterization of Parvovirus in Ducks in Northern Vietnam Reveal Evidence of Recombination

Hieu Van Dong, Giang Thi Huong Tran, Huong Thi Thu Nguyen, Tuong Manh Nguyen, Dai Quang Trinh, Van Phan Le, Kiattawee Choowongkomon, and Jatuporn Rattanasrisomporn

1. Introduction

Waterfowl parvoviruses are highly contagious, lethal pathogens for goslings and ducklings. Waterfowl parvoviruses belong to the species *Anseriform dependoparvovirus 1* of the genus *Dependoparvovirus* within the *Paroviridae* family. The genus *Dependoparvovirus* can be divided into two divergent groups, consisting of Muscovy duck parvovirus (MDPV)-related groups and goose parvovirus (GPV)-related groups [1]. GPV, the agent of Derzy’s disease, causes the disease in young geese and Muscovy ducks with mortality rates up to 90% [2], while MDPV causes high mortality, watery diarrhea, wheezing, locomotor
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dysfunction, and stunning in Muscovy ducks [3]. Both GPV and MDPV can show 70–100% morbidity and mortality during the first 3–4 weeks of age.

Waterfowl parvovirus is characterized as a non-enveloped virus. The viral genome consists of a single-strand DNA with around 5100 bp. The translated regions contain two open reading frames (ORFs). The left ORF encodes for the non-structural protein, while the remaining ORF encodes for capsid proteins VP1, VP2, and VP3 [4,5]. Both GPV and MDPV are antigenically related to each other as they share about 85% protein sequence homology [6]. In detail, they show nucleotide differences in VP1 of about 20–24%, whereas these differences in VP1 within the GPV and MDPV groups are only 0.1–7% and 0.1–1.9%, respectively [6–9].

It was reported that recombination events occurred among paroviruses. As a result, a novel strain may be generated from the recombination event between vaccine and wild-type strains [10]. In 2015, Chen et al. found a novel goose parvovirus (NGPV) causing a severe disease in duck flocks in China [11]. Later, other studies noted that NGPV strains induced short beak and dwarfism syndrome (SBDS) in China, Egypt, and Poland [12–14]. NGPV viruses were considered to derive from classical GPV strains. Furthermore, a novel recombinant MDPV (rMDPV) was reported in ducks from several provinces in mainland China, where it caused high mortality and embolism in the intestinal tracts of infected ducklings aged less than 3 weeks [15–17].

In Vietnam in 2019, waterfowl parvoviruses have been reported in sick ducks with short beak and dwarfism syndrome. The detected waterfowl parvovirus was genetically grouped with the NGPV group based on the partial NS and VP1 genes [18]. However, information on waterfowl parvoviruses has been limited up to now. Therefore, we investigated infections of waterfowl parvoviruses in ducks farmed in some provinces of northern Vietnam. The current study also carried out molecular characterization of the near-complete waterfowl parvovirus genome.

2. Materials and Methods

2.1. Ethics Statement

This research did not contain any studies involving human participants. Collection of duck tissue samples was conducted by the Vietnam National University of Agriculture after receiving institutional approval and permission from the owners of the ducks.

2.2. Samples

Tissue samples (brain, lung, liver, kidney, and Fabricius) from 130 broiler ducks aged 2–7 weeks were obtained from Thainguyen, Bacgiang, Haiduong, Thaibinh, Hungyen, and Hanoi in 2021 (Figure 1) and used in this study. From each flock, two to six diseased ducks were collected by the local veterinarian and transferred to the Vietnam National University of Agriculture for further analysis. The parvovirus vaccine was not used in tested duck farms. From each duck, collection of pooled tissue samples was performed. A 10% homogenate was prepared in phosphate-buffered saline.

2.3. DNA Extraction and Polymerase Chain Reaction (PCR)

DNA was extracted from the homogenized samples using Viral Gene-spin™ Viral DNA/RNA Extraction Kits (iNtRON Biotechnology, Seoul, Korea) according to the manufacturer’s instruction. Primers, PV-F, and PV-R were used to amplify the target 537 bp of the partial replication (Rep) gene or the parvovirus genome identification (Table 1), as previously described [4]. The thermal conditions were: 95 °C for 5 min, 35 cycles of 95 °C for 30 s, 55 °C for 30 s, 72 °C for 30 s, and extended at 72 °C for 10 min. A 1.5% agarose gel was used to run the PCR product. The product was observed by UV light.
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Table 1. Primers used in this study.

| Target | Name    | Sequence (5'–3')       | PCR Product (bp) | Location of Target Genes on the Viral Genome | Reference |
|--------|---------|------------------------|------------------|---------------------------------------------|-----------|
| PCR    | PV-F    | CCAAGCTACAAACAACCACAT  | 539              |                                              | [4]       |
|        | PV-R    | TGAAGCATACATGCTATGGAAGG|                  |                                              |           |
|        | GPV-P1-F| CTTATTGAGGGGTTGCTGT    | 176              | 5'-UTR                                      | [14]      |
|        | GPV-P1-R| GCATCGCGCTGCTCAACCTAA  |                  |                                              |           |
|        | GPV-P2-F| GCATCGCGCTGCTCAACCTAA  | 1033             | 5'-UTR/Rep                                  | [14]      |
|        | GPV-P2-R| AATTCAATGAGCAAATCAACAAAG|                |                                              |           |
|        | GPV-P3-F| GCCTTTATTTAATCTGGCT    | 1443             | Rep                                         | [14]      |
|        | GPV-P3-R| GCTTTTACATTCGCCCAC     |                  |                                              |           |
|        | GPV-P4-F| CTTGATGATGCTGAAAAATGAAC|                | Rep/Cap                                    | [14]      |
|        | GPV-P4-R| GCCATGCTGCTCAACCTAAGC  | 1446             | Cap/3'-UTR                                 | [14]      |
|        | GPV-P5-F| GCACCTACACGAGACTTAGACA  | 1170             | Cap/3'-UTR                                 | [14]      |
|        | GPV-P5-R| GCATCGCGCTGCTCAACCTAA  |                  |                                              |           |

2.4. Nucleotide Sequencing and Phylogenetic Analyses

Five pairs of primers, GPV-P1-F/R, GPV-P2-F/R, GPV-P3-F/R, GPV-P4-F/R, and GPV-P5-F/R (Table 1), were used to amplify the near-whole genome sequence of the waterfowl parvovirus strains, as described previously [14]. Electrophoresis was performed with 1.5% agarose gel to separate the PCR products. Regarding the purification of PCR products, GeneClean® II Kits (MP Biomedicals, Santa Ana, CA, USA) were used. Sequencing of the waterfowl parvovirus strains was demonstrated by 1st BASE, Malaysia.

The Clustal W multiple alignment tool [19] in BioEdit v.7.2.5 [20] was used to align and analyze the nucleotide sequences and deduced aa sequences derived from GPV. Homology in nucleotide and aa sequences was examined using the GENETYX v.10 software (GENETYX Corp., Tokyo, Japan) and compared with other publicly available sequences using the BLAST program. A maximum likelihood method with the Hasegawa–Kishino–Yano model of nucleotide substitutions was used to construct the phylogenetic tree based on nucleotide sequences of 4 currently identified Vietnamese and 40 foreign waterfowl parvovirus strains (Table 2). The confidence values on phylogenetic trees were assessed based on bootstrapping with 1000 replicates using the MEGA6 6.06 version [21]. The near-whole genomes of the current waterfowl parvovirus strains were submitted to GenBank with accession numbers OP265005–OP265008.
Table 2. Description of waterfowl parvovirus isolates compared in this study.

| GenBank Accession No. | Strain | Location | Source | Year |
|-----------------------|--------|----------|--------|------|
| MN549533.1            | GDQY1802 | China    | Duck   | 2018 |
| MN549532.1            | GDSC1902 | China    | Duck   | 2018 |
| KT935531.2            | JS1     | China    | Duck   | 2015 |
| KX384726.2            | DS15    | China    | Duck   | 2015 |
| KU641558.1            | CVS01   | China    | Duck   | 2015 |
| KY679174.1            | SC16    | China    | Duck   | 2016 |
| MF441227.1            | AH1605  | China    | Duck   | 2016 |
| MH444513.1            | AH      | China    | Duck   | 2018 |
| MN415972.1            | SD1228  | China    | Duck   | 2018 |
| MN415972.1            | SD1228  | China    | Duck   | 2018 |
| MN356044.1            | SDJN19  | China    | Duck   | 2019 |
| EU833392.1            | VG32/1 vaccine | Taiwan | Goose | 2008 |
| EU833389.1            | 82-032IV vaccine | Taiwan | Goose | 1982 |
| MT646164.1            | AHAU30  | China    | Duck   | 2019 |
| EF515837.1            | DY      | China Muscovy duck | 2007 |
| MT646163.1            | AHAU41  | China    | Duck   | 2019 |
| MF942876.1            | SQ0412  | China    | Goose  | 2017 |
| KY475362.1            | RC16    | China    | Goose  | 2016 |
| KC996730.1            | YZ99-6  | China    | Goose  | 1999 |
| KM272560.1            | LH      | China    | Goose  | 2012 |
| KR136258.1            | Yan-2   | China    | Goose  | 2013 |
| KR091960.1            | YZ      | China    | Goose  | 2013 |
| EU833391.1            | 06-0329 | Taiwan   | Goose  | 2008 |
| KC1841333.1           | E       | China    | Goose  | 2021 |
| MW386077.1            | Corum/19 | Turkey | Goose  | 2019 |
| MW386079.1            | Yozgat/19 | Turkey | Goose  | 2019 |
| MW386078.1            | Konya/19 | Turkey | Goose  | 2019 |
| MF441223.1            | SDHZ1604 | China   | Duck   | 2016 |
| MF441221.1            | SDLY1512 | China   | Duck   | 2015 |
| MF441226.1            | JS1603  | China    | Duck   | 2016 |
| MF441222.1            | SDLY1602 | China   | Duck   | 2016 |
| MH444514.1            | GD      | China    | Duck   | 2016 |
| MK000549.1            | HuN001  | China    | Duck   | 2018 |
| MK737642.1            | HN1P    | China    | Duck   | 2019 |
| KY511124.1            | SD      | China    | Duck   | 2015 |
| KT3432533.1           | sdlick1 | China    | Duck   | 2015 |
| KM093740.1            | MDPV-GX5 | China Muscovy duck | 2011 |
| KU844282.1            | P1      | China    | Muscovy duck | 2016 |
| KT865605.1            | FZ91-30 vaccine | China Muscovy duck | 1991 |
| KU844281.1            | P       | China    | Muscovy duck | 1988 |

2.5. Analyses of Recombination and Natural Selection Profiles

The current and other waterfowl parvovirus sequences available in GenBank were determined to be recombination events using the Recombination Detection Program (RDP) version Beta 4.97 [22] (including RDP, GENECONV, BootScan, MaxChi, Chimaera, SiScan, Phyl-Pro, LARD, and 3Seq methods) with default settings. An algorithm producing a result that had p value lower 0.05 was regarded as dependable. The identification of sites under positive or negative selection was evaluated by following a FUBAR (a Fast Unconstrained Bayesian AppRoximation) method (accessed on 1 July 2022) [23].

2.6. Statistical Analysis

Fisher’s exact test was used to identify significant differences in the rate of waterfowl parvovirus genome detection between geographical regions, age, or flock-size groups. A value of p < 0.05 was considered statistically significant.

3. Results

3.1. Detection of Waterfowl Parvovirus Genome in Field Samples

Of the 130 samples tested, 26 (20%) were positive for the waterfowl parvovirus genome based on the PCR method. Among the provinces/cities, Haiduong had the highest rate (33.33%), followed by (p > 0.05) Thainguyen (26.67%), Hanoi (22.92%), and Hungyen (20%), that were all significantly higher (p < 0.05) than that of Thaibinh (14.29%) and Bacgiang
(13.33%) (Table 3). Of the 38 farms tested, 14 were positive for the waterfowl parvovirus genome (Table 3).

Table 3. Detection of waterfowl parvovirus genome in field duck samples obtained from different northern provinces/cities of Vietnam.

| Province/City | No. of Collected Samples | Gene-Positive Samples/(%) | No. of Flocks | Gene-Positive Flocks/(%) |
|--------------|--------------------------|---------------------------|---------------|-------------------------|
| Hanoi        | 48                       | 11/(22.92) a,b            | 11            | 4/(36.36) c            |
| Haiphong     | 6                        | 2/(33.33) a,b             | 2             | 1/(50.00) c,d          |
| Thua Thien   | 15                       | 4/(26.67) a,b             | 3             | 2/(66.67) d            |
| Bac Giang    | 30                       | 4/(13.33) b               | 11            | 3/(27.27) c,e          |
| Ha Tinh      | 21                       | 3/(14.29) b               | 7             | 2/(28.57) c,e          |
| Hung Yen     | 10                       | 2/(20.00) a,b             | 4             | 2/(50) c,d            |
| Total        | 130                      | 26/(20.00)                | 38            | 14/(36.84)             |

Letters a,b,c,d,e indicate that the groups were significantly (p < 0.05) different from each other.

The rate of parvovirus-positive ducks at age 2–4 weeks was 37.04% (10/27), which was highly significantly greater those aged <2 weeks (9.09%; p < 0.01) and >4 weeks (16.30%; p = 0.01) (Table 4). The flock levels were categorized as level 1 (number of ducks lower than 500, small scale), level 2 (number of ducks ranging 500–1000, medium scale), or level 3 (number of ducks more than 1000, large scale). The rates of parvovirus detection for the level 1 and level 3 flocks were 31.03% and 29.73%, respectively, which were significant (p < 0.05) higher than for level 2 flocks (9.38%), as shown in Table 4.

Table 4. Detection of waterfowl parvovirus genome in field duck samples by age and flock size.

| Criteria | Ages/Duck Heads | Samples (n) | Gene-Positive Samples/(%) |
|----------|-----------------|-------------|--------------------------|
| Age (weeks) |                |             |                          |
| <2       | 11              | 1/(9.09) a  |                          |
| 2–4      | 27              | 10/(37.04) b|                          |
| >4       | 92              | 15/(16.30) a|                          |
| <500     | 29              | 9/(31.03) a |                          |
| 500–1000 | 64              | 6/(9.38) b  |                          |
| >1000    | 37              | 11/(29.73) a|                          |

Letters a,b indicate that the groups were significantly (p < 0.05) different from each other.

3.2. Characterization of Waterfowl Parvovirus Genome and Protein

The near-complete genome of the four Vietnamese waterfowl parvoviruses obtained in this study was 4750 nucleotides in length. The viral genomes of the four Vietnamese waterfowl parvovirus types consisted of two ORFs. Neither deletion nor insertion mutations were detected in the translated region of the current parvovirus.

Regarding genetic analysis, four parvovirus-detected samples obtained at different locations were randomly determined for viral genome sequencing. The alignment and comparison of the near-whole genome (4750 bp) of the four current strains and other reported sequences from GenBank were performed. The nucleotide identity ranged from 99.29% to 99.87% among the four Vietnamese waterfowl parvovirus strains obtained in this study. Among these, the highest nucleotide identity was between Vietnam/VNUA-07/2021 and Vietnam/VNUA-26/2021 (99.87%) while the lowest was between Vietnam/VNUA-30/2021 and Vietnam/VNUA-94/2021 (99.29%). Comparing the viral genomes from the four Vietnamese parvovirus strains in this study and those sequences abroad, four Vietnamese strains shared nucleotide identity of 99.17% (Vietnam/VNUA-94/2021 versus China/HuN001), 99.04% (Vietnam/VNUA-07/2021 versus China/HuN001, Vietnam/VNUA-26/2021 versus China/HuN001), and 98.82% (Vietnam/VNUA-30/2021 versus China/HuN001).

Phylogenetic analysis based on the near-complete genome (4880 bp), full-length Cap (2199 bp), and full-length Rep (1884 bp) gene sequences indicated that the current Vietnamese parvovirus strains belonged to the NGPV group. The four Vietnamese parvovirus strains obtained differed from vaccine strains Taiwan/VG32/1/2008 (EU583392.1), Taiwan/82-
The present Vietnamese parvovirus strains were clustered with those of viral strains from China (Figures 2–4).

Figure 2. Phylogenetic trees of the near-complete genome (4750 bp) sequences of Vietnamese parvovirus strains compared with those available in GenBank. GenBank sequences are indicated by the country name/accession number. The maximum likelihood method in the MEGA6 software was used to establish phylogenetic trees (1000 bootstrap replicates). Numbers at each branch point indicate bootstrap values $\geq 50\%$ in the bootstrap interior branch test. The current Vietnamese and vaccine strains are indicated by circles and diamonds, respectively.
Figure 3. Phylogenetic trees of the full-length Cap gene (2199 bp) sequences of Vietnamese parvovirus strains compared with those available in GenBank. GenBank sequences are indicated by the country name/accession number. The maximum likelihood method in the MEGA6 software was used to establish phylogenetic trees (1000 bootstrap replicates). Numbers at each branch point indicate bootstrap values $\geq 50\%$ in the bootstrap interior branch test. The current Vietnamese and vaccine strains are indicated by circles and diamonds, respectively.
Deduced aa sequences of the Rep and Cap proteins of the Vietnamese NGPV strains were compared with other waterfowl strains in various lineages. Four and fourteen major variable amino acid substitutions were detected on Rep and Cap proteins, respectively, of the Vietnamese NGPV strains in the current study, compared to other GPV and MDPV viruses (Tables 5 and 6).
Table 5. Amino acid substitutions in Rep protein of Vietnamese NGPV strains.

| Virus Strain/Consensus | Sites in Rep Protein |
|------------------------|----------------------|
|                        | 154 | 551 | 555 | 560 |
| GPV                    | T   | R   | N   | C   |
| MDPV                   | T   | K   | D   | C   |
| Vietnamese NGPVs      | T/S | K/R | P/T | G/C |

* Deduced amino acid consensus was released with 102 waterfowl parvovirus sequences retrieved from GenBank by using GENETYX version 10.

Table 6. Amino acid substitutions in Cap protein of Vietnamese NGPV strains.

| Virus Strain/Consensus | Sites in Cap Protein |
|------------------------|----------------------|
|                        | 142 | 144 | 178 | 180 | 206 | 386 | 390 | 448 | 449 | 451 | 458 | 461 | 507 | 509 |
| GPV                    | D   | V   | S   | A   | A   | N   | A   | D   | G   | R   | A   | G   | D   | Q/E |
| MDPV                   | E   | V   | N   | G   | A   | N   | A   | D/N | S/G | R   | A   | G   | D   | E   |
| Vietnamese NGPVs      | E/D | I/V | T/S | A/V | T/A | N/D | A/P | D/G | S/R | R/G | A/T | G/R | E/D/A | Q/P |

* Deduced amino acid consensus was released with 102 waterfowl parvovirus sequences retrieved from GenBank by using GENETYX version 10.

3.3. Recombination Analysis

Recombination analyses of the full-length Cap gene sequences suggested that Vietnam/VNUA-26/2021 resulted from a recombination event. The major and minor parents were the Duck/China/GXN45/2017 NGPV and Goose/China/YZ/2013 GPV viruses. This putative recombination event was detected by seven out of the nine methods using RDP 4 (Table 7). The two breakpoints were detected and located at residues 112 and 620 of the putative recombination event was detected by seven out of the nine methods using RDP 4 (Table 7). The two breakpoints were detected and located at residues 112 and 620 of the putative recombination event. The major and minor parents were the Duck/China/GXN45/2017 NGPV and Goose/China/YZ/2013 GPV viruses. This putative recombination event was detected by seven out of the nine methods using RDP 4 (Table 7). The two breakpoints were detected and located at residues 112 and 620 of the putative recombination event. The major and minor parents were the Duck/China/GXN45/2017 NGPV and Goose/China/YZ/2013 GPV viruses. This putative recombination event was detected by seven out of the nine methods using RDP 4 (Table 7). The two breakpoints were detected and located at residues 112 and 620 of the putative recombination event. The major and minor parents were the Duck/China/GXN45/2017 NGPV and Goose/China/YZ/2013 GPV viruses. This putative recombination event was detected by seven out of the nine methods using RDP 4 (Table 7). The two breakpoints were detected and located at residues 112 and 620 of the putative recombination event.

Table 7. Recombination statistics of Vietnam/VNUA-26/2021 Cap gene sequence using RDP 4.

| Method          | Recombination p-Value |
|-----------------|-----------------------|
| RDP             | 1.89 × 10⁻²           |
| GENECONV        | 4.77 × 10⁻⁴           |
| BootScan        | 2.84 × 10⁻²           |
| MaxChi          | 2.72 × 10⁻⁵           |
| Chimaera        | 8.11 × 10⁻⁶           |
| SiScan          | 5.41 × 10⁻¹²          |
| PhyIPro         | -                     |
| LARD            | -                     |
| SSSeq           | 4.48 × 10⁻³           |

Recombination events with p-value < 0.05 were regarded as reliable.

Figure 5. Detection of recombination events using BootScan analysis of full-length Cap gene sequences of Vietnam/VNUA-26/2021 NGPV strains. The pairwise distance model with window size 200, step size 20, and 1000 bootstrap replicates was generated by the RDP 4 program.
3.4. Evolutionary Analysis of Viral Genome

Analyses of the natural selection profiles of the Vietnamese NGPV sequences indicated that eight residue sites (137, 147, 157, 159, 161, 168, 472, and 686) of the Cap protein and one residue (508) site of the Rep protein were from negative selection. However, only one positive selection was found at the residue 507 site of the Cap protein (Table 8).

Table 8. Natural selection profile in waterfowl parvovirus capsid and replication protein of four current Vietnamese NGPV strains.

| Protein | Site | a    | b    | b–a  | Prob [a > b] | Prob [a < b] | Bayes Factor [a < b] |
|---------|------|------|------|------|--------------|--------------|---------------------|
| Capsid  | 137  | 26.755 | 2.394 | −24.361 | 0.912 | 0.063 | 0.083 |
|         | 147  | 26.312 | 2.169 | −24.142 | 0.916 | 0.06 | 0.078 |
|         | 157  | 25.594 | 2.021 | −23.573 | 0.915 | 0.061 | 0.079 |
|         | 159  | 26.745 | 2.188 | −24.557 | 0.917 | 0.06 | 0.079 |
|         | 161  | 24.335 | 2.109 | −22.226 | 0.906 | 0.068 | 0.089 |
|         | 168  | 25.174 | 2.226 | −22.948 | 0.908 | 0.065 | 0.086 |
|         | 472  | 24.338 | 2.108 | −22.23  | 0.906 | 0.068 | 0.089 |
|         | 507  | 4.423  | 33.746 | 29.323 | 0.034 | 0.926 | 15.48 |
|         | 686  | 25.729 | 2.158 | −23.571 | 0.912 | 0.063 | 0.083 |
| Replication | 508 | 30.806 | 3.718 | −27.088 | 0.905 | 0.062 | 0.077 |

4. Discussion

Understanding waterfowl parvovirus infections is important as they are associated with SBDS in ducks with a high or low mortality [12–14]. Recently, waterfowl paroviruses were characterized based on partial sequences of the VP1 and NS1 gene sequences and divided into the NGPV group in Hungyen province [18]. The Vietnamese waterfowl parovirus genome sequences are still limited in GenBank; additional sequence data of the complete genome are necessary to further characterize and understand the evolution of viral strains. This was the first study to conduct epidemiological analysis and to further characterize the whole-genome sequences (4880 bp) of Vietnamese waterfowl parovirus strains.

In the current study, the waterfowl parovirus genome was detected in 20% of duck samples, which was lower than that of ducklings from outbreak farms with SBDS in tissue samples (33/33 (100%)) or with no clinical signs (7/12 (58.33%)) [24]. The differences in the sampling numbers, location, time, and sensitivity of the detection methods were considered a reasonable cause for the variation in the result. Furthermore, 36.84% of the duck farms were positive for the viral genome in northern provinces in Vietnam. The current results suggested that waterfowl parovirus may circulate and affect duck production in Vietnam. In the current study, we first found a moderate waterfowl parovirus genome frequency of detection in Vietnamese ducks at age 2–4 weeks (37.04%), with the frequency of detection of viral genome higher than that in younger ducks aged <2 weeks (9.09%) and older ducks aged >4 weeks (16.30%). These findings suggested that during age 2–4 weeks the ducklings may be susceptible to viral infection.

Genetic and phylogenetic analyses indicated that the Vietnamese and Chinese strains were clustered in a single genogroup belonging to NGPV (Figures 2–4). The results suggested that these viruses may have a similar origin, or that these viruses were widespread in Asia. In addition, the classification of the Vietnamese NGPV sequences was similar based on the complete genome, Rep gene, and Cap gene sequences, suggesting that using each gene sequence could characterize the viral strains. Amino acid substitutions on Rep and Cap proteins play critical roles in the change of host range and pathogenicity. In terms of Rep protein, a previous study reported that substitution at residue 140 (A to S), which was not be found in the current four Vietnamese strains, was associated with viral transcription [25]. Yu et al. noted that five substitutions at residues 494, 553, 555, 573, and 594 on Rep protein play roles in the antigenic epitopes of B cells [26]. Only one substitution at residue 555 N/D to P/T was detected in the present study. The roles of the remaining
three substitutions on Rep protein of the Vietnamese strains were still unclear. In this study, 14 substitutions on Cap protein did not match with residues 35 and 660, which may be related to the host range identification of NGPV [27].

SBDS was recognized in the NGPV group in China in 2015 [11]. Since then, this pathogen has been critical in causing disease in ducks, mainly in China but also in other countries. In the current study, the four NGPVs were detected based on their complete genome sequences. The four Vietnamese NGPVs obtained were closely related to NGPV in China in 2019 (HuN18). The HuN18 NGPV virus has been isolated from ducks with BADS in China [28]. Therefore, it could be speculated that the current Vietnamese NGPV strains may be high-virulence strains.

Recombination has been reported in the evolutionary processes of waterfowl parvoviruses and described [17,28–30] based on the complete genome sequences. A recombination event may generate a novel waterfowl parvovirus strain [10]. The current study revealed the first evidence supporting a recombination event in the Vietnamese NGPV strain in the protein-coding region of the Capsid gene. In general, recombination usually occurs between viruses that are located closely to each other. However, the major and minor parents were from China. The explanation for this situation is that Vietnam and China share thousands of kilometers along their border. The recombination event could have occurred elsewhere in China and then could have been transferred to Vietnam through daily trading of live birds along the border between Vietnam and China [31]. Recombination may occur between the Vietnamese waterfowl parvovirus strains, which might be detected if there are sufficient data. Analysis of the additional viral genomes should be further studied.

Another evolutionary process is natural selection. Fan et al. [32] reported nine sites on VP protein that were estimated as positively selected sites, that may have been associated with host range [32]. In the current study, only one positive selection on a VP protein was found (residue 507), which has not been characterized. Further studies should be conducted to clarify this point.

5. Conclusions

A high positive rate of waterfowl parvovirus was detected among young ducks, aged 3–4 weeks, in flocks in northern Vietnam. The current study revealed that the four Vietnamese waterfowl parvovirus strains belonged to the NGPV group, based on phylogenetic and molecular analyses of the whole genome. The characterization of NGPV strains circulating in Vietnam was first reported based on viral whole-genome sequences. This study also detected a recombination event in the Vietnamese NGPV strain. Only one aa sequence on a Cap protein was identified under positive selection.

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