

Molecular Characteristics and Phylogenetic Analysis of Short Beak and Dwarfism Syndrome-Related Novel Goose Parvovirus

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Research Article

Keywords: short beak and dwarfism syndrome, goose parvovirus, molecular characteristics, medium egg lethal dose, inverted terminal repeats, phylogenetic tree

DOI: <https://doi.org/10.21203/rs.3.rs-349163/v1>

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Abstract

Short beak and dwarfism syndrome (SBDS) emerged in cherry valley duck flocks in China in 2015, and novel goose parvovirus (NGPV) was proved to be the etiological agent of SBDS. To date, whether SBDS-related NGPV isolates possess common molecular characteristics remains unknown. In this study, three new NGPV strains (namely, SDHT16, SDJN19, and SDLC19) were isolated from diseased ducks showing typical SBDS and successfully passaged in embryonated goose or cherry valley duck embryo. The whole genomes of three NGPV strains shared 98.9%–99.7% homologies between each other but showed slightly lower homologies (95.2%–96.1%) with the classical GPV strains. A total of 16 common amino acid point mutations were produced in the VP1 proteins of six NGPV strains (SDHT16, SDJN19, SDLC19, QH, JS1, and SDLC01) compared with the classical Chinese GPV strains, among which nine amino acid sites were identical to the European GPV strain B. The non-structural protein Rep1 of the six NGPV strains generated 12 common amino acid mutations compared with the classical GPV strains. The phylogenetic analysis indicated that the Chinese NGPV strains clustered with the European SBDS-related NGPV strains, forming a separate branch, distinct from the group formed by the classical GPV strains. Taken together, the present study unveils the common molecular characteristics of the NGPV isolates and directs the conclusion that the Chinese NGPV isolates probably originate from a common ancestor with the European SBDS-related NGPV.

Introduction

Short beak and dwarfism syndrome (SBDS) emerged in cherry valley duck flocks in China in 2015 with the morbidity rate ranging from 5–25% [1, 2, 11]. Ducks with this disease are characterized by a short beak, protruding tone, fragile bone, and retarded growth [2, 12]. The poor body weight gain and heterogeneity within the flock lead to economic losses to the duck meat industry. SBDS was first reported in the 1970s in France in mule ducks and also reported in Poland in 1995 [14].

The causative agent of SBDS is closely related to the classical goose parvovirus (GPV), tentatively named as novel GPV or GPV variant [20]. GPV and Muscovy duck parvovirus are classified in the genus of *Dependoparvovirus* in the *Parvoviridae* family [3]. The viral genome has a ~ 5.1-kb single-stranded linear DNA with equal polarities encapsidated in the icosahedral capsid. The genome is flanked with identical inverted terminal repeats (ITRs) and contains two ORFs, the left encoding the Rep protein and the right encoding the capsid protein [21]. The Rep ORF produces the largest protein Rep1 and several low-molecular-weight Rep proteins from spliced mRNAs, which are involved in genome replication, interaction with ITR, and modulation of the downstream P41 promoter [10, 13, 16, 18]. By alternative mRNA splicing and protease cleavage, the right ORF produces three structural proteins (VP1, VP2, and VP3), which assemble the viral capsid with a ratio of 1:1:8 [7, 16].

The GPV DNA can be detected by PCR from the internal organs of diseased ducks showing SBDS; however, viral isolation from tissues (liver, spleen, and kidney) is not always successful [1, 20]. Viral isolation can be successfully achieved in susceptible duck embryos through the inoculation of tissue

homogenate. Blind passages in embryonated duck eggs are necessary to establish good adaptation for the primary novel goose parvovirus (NGPV) isolates [1]. Since the outbreak of SBDS in China, several genome sequences of NGPV isolates have been sequenced [1, 12, 20], but whether these isolates possess common molecular characteristics and similar propagation capability in avian embryos remains unclear. Moreover, the evolution correlation between the NGPV isolates and the classical GPV strains and the European SBDS-related GPV isolates remains to be elucidated.

In the present study, three NGPV strains were isolated from SBDS cases during disease outbreaks in cherry valley duck flocks between 2016 and 2019, and their genomes were sequenced. Sequence alignments revealed that all Chinese NGPV strains harbored common amino acid mutations in the coding proteins compared with the classical GPV strains. Moreover, two NGPV strains isolated in different years exhibited obviously differential propagation ability in embryonated duck embryos. Phylogenetic analysis indicated that the Chinese NGPV isolates probably originated from the European SBDS-related NGPV.

Materials And Methods

Viral isolation and passage

For the diseased cherry valley ducks showing typical SBDS, their internal organ tissues, such as liver, kidney, and spleen, were sampled between 2016 and 2019 in Shandong Province, China. The tissue samples were placed in sterile saline for homogenization. The homogenate was clarified by centrifugation at $3000 \times g$, and the supernatant was transferred and passed through a filter with a $0.22\text{-}\mu\text{m}$ pore diameter. The filtrate was added with penicillin (1000 IU/mL) and streptomycin (1000 $\mu\text{g}/\text{mL}$) and then used to inoculate susceptible 9-day-old embryonated goose eggs via the chorioallantoic membrane route. The eggs were incubated at $38\text{ }^{\circ}\text{C}$ for 12 days and candled three times every day. The dead embryos were placed at $4\text{ }^{\circ}\text{C}$ for 6 h, and then the allantoic fluid was pooled. The embryos were further examined for pathogenic lesions. The first-generation allantoic fluid containing virus was 1:10 diluted with sterile saline, and the dilution was used to inoculate another five 9-day-old goose eggs or cherry valley duck eggs for passage.

Determination of medium egg lethal dose

To evaluate whether the NGPV isolates isolated in 2016 and 2019 possess differential propagation capability in embryonated duck eggs, the medium embryo lethal dose (ELD_{50}) against two NGPV strains was calculated in accordance with the method of Reed and Muench [17]. Briefly, the viral stock in the form of allantoic fluid was diluted in sterile saline to give a 10-fold dilution series between 10^{-1} and 10^{-7} . For each dilution, 0.2 mL was inoculated into 9-day-old susceptible cherry valley duck eggs (five eggs every dilution) via the allantoic cavity route. The inoculated eggs were continuously incubated at $37\text{ }^{\circ}\text{C}$ and examined three times daily for seven days. The times of any embryo deaths were recorded by hours. The ELD_{50} experiments were repeated for three times and the mean value was calculated.

Whole genome sequencing

Seven sets of primers were designed on the basis of the genomic sequence of the classical GPV strain LH (accession number KM272560, Table 1). These primers were used to amplify the overlapping fragments by using PrimeSTAR Max DNA polymerase (Takara, Dalian, China). Except for the 200-bp terminal fragments containing half of the ITR, five internal DNA fragments were gel-purified using the TIANGel Midi Purification Kit (Tiangen, Peiking, China) and then subjected to direct sequencing using the amplification primers. An “A” tail was added to the 200-bp terminal fragments using the Taq DNA polymerase. The fragments were then ligated with the pMD19T vector (Takara, Dalian, China) and transformed into competent DH5 α cells. Positive clones were screened for sequencing. The obtained sequences were combined to assemble the whole genome sequence using the SeqMan program packaged in Lasergene 7.0 (DNASTAR, Madison, USA).

In addition, the ITR nucleotide sequences of three strains were confirmed by sequence analysis of recombinant plasmids carrying the viral DNAs purified directly from the virions without PCR amplification. Virus purification and concentration from the allantoic fluid and extraction of genomic DNA were performed as previously described [19]. The extracted single-stranded DNA (ssDNA) was suspended in STE buffer (10 mM Tris, 1 mM EDTA, 100 mM NaCl, pH 8.0) and annealed by heating to 95 °C for 5 min followed by slow cooling to 55 °C. The double-stranded DNA (dsDNA) was digested with *Nco*I, separated by electrophoresis, and purified using a gel extraction kit (Tiangen, Peking, China). The 1.3-kb DNA fragment containing the 3'-end ITR was ligated into the *Nco*I-*Sma*I site of the pBSKNB plasmid [19] and used to transform competent cells of the Sure strain of *E. coli*. The positive clones were chosen for sequencing. To overcome difficulties in sequencing ITR regions with hairpin structure, the ITR was digested by *Sph*I, which cut the ITR in the middle loop region, and the resulting fragments were subcloned into the pUC18 plasmid for sequencing.

Sequence alignment and phylogenetic analysis

Fourteen strains, including six classical GPV strains and eight SBDS-related NGPV strains, were used in this study for sequence alignment and phylogenetic tree construction. JS1, QH15, and SDLC01 were three representative NGPV strains that were isolated at the early stage of the SBDS outbreak in China (Table 2). Except for D146/02 and D697/3/06, both of which are European SBDS-related NGPV isolates (Palya *et al.*, 2009) and only have partial VP1 gene sequences deposited in GenBank, the 12 other strains have complete genome sequences available in GenBank. Homology comparisons of the coding proteins among the classical GPV strains and SBDS-related NGPV strains were performed using the MegAlign program packaged in Lasergene 7.0. Given that the European SBDS-related NGPV strains only have partial VP1 gene sequences (at the N terminus) available in GenBank, the phylogenetic tree was constructed on the basis of a 427-bp VP1 gene fragment to explore the possible evolutionary origin of the Chinese SBDS-related NGPV strains. The trees were constructed using MEGA 7.0 [9] (Kumar *et al.*, 2016) by the maximum likelihood model and bootstrap values were calculated based on 1000 replicates.

Statistical analysis

The two-tailed student's t-test was adopted to compare the mean death time (MDT) of duck embryos post-inoculation between the SDHT16 and SDJN19 isolate by using GraphPad Prism software version 6.01 (GraphPad Software Inc., San Diego, CA, USA) . $P < 0.05$ was considered statistically significant. The data were expressed as mean \pm standard deviation (SD).

Results

Viral isolation and passage in embryonated goose eggs

Three organ tissue samples were collected from diseased cherry valley ducks showing typical SBDS between 2016 and 2019 in Shandong Province, China. The three samples were positive in PCR detection using specific GPV primers. Each sample was used to inoculate five susceptible 9-day-old embryonated goose eggs. The result showed that 1, 2, and 1 of five embryonated goose eggs died between 234 h and 288 h post-inoculation. The three viral isolates were successfully passaged in another five 9-day-old embryonated goose eggs and killed all the goose embryos in a shorter time (between 112 h and 190 h) post-inoculation. Hemorrhagic lesions in the head, neck, and legs typical of parvoviral infection were observed (Fig. 1). The three NGPV isolates were named SDHT16, SDJN19, and SDLC19.

NGPV propagation in cherry valley duck embryos and ELD₅₀ calculation

Strain SDHT16 and SDJN19 were successfully passaged in 9-day-old embryonated cherry valley duck eggs (Fig. 1). Strain SDHT16 killed 16 duck embryos between 106 h and 197 h (averaging 150 h) post-infection, and strain SDJN19 killed 16 duck embryos between 69 h and 163 h (averaging 104.6 h). Statistical analysis indicated that the mean death time between the two NGPV strains was significantly different ($P < 0.05$) (Fig. 2).

Furthermore, the calculation of the medium lethal egg doses (ELD₅₀) was performed in embryonated cherry valley duck eggs. The ELD₅₀ values of strain SDHT16 and SDJN19 reached $5 \times 10^{2.5}/\text{mL}$ and $5 \times 10^{5.4}/\text{mL}$, respectively, demonstrating that strain SDJN19 has a better propagation capability in embryonated cherry valley duck eggs than strain SDHT16.

Genome sequencing and homology comparison

The whole genomes of three NGPV isolates were amplified by PCR and sequenced. The genome of strain SDHT16 is 5050 bases long, which is four bases shorter than that of strain SDJN19 and SDLC19. Nucleotide deletions or insertions occurred in ITR with no exception. The three NGPV strains shared 98.9%–99.7% similarities at the genomic level but displayed slightly lower nucleotide homologies (95.2%–96.1%) with the classical GPV strains (strain B, 82-0321, 06-0329, YZ99-6, and LH). The genome sequences of three NGPV strains were deposited in GenBank with accession numbers MN356043, MN356044, and MN356045.

Sequence comparison of the Rep1 proteins

The Rep1 gene of the three NGPV strains (SDHT16, SDJN19, and SDLC19) was composed of 1884 nucleotides, coding 627 amino acids. Homology comparison indicated that the Rep1 gene of three NGPV strains shared 99.3%–99.7% similarities between each other but displayed 96.0%–96.5% similarities with the Chinese GPV strains (82-0321, 06-0329, YZ99-6, and LH) and 96.8%~96.9% similarities with the European GPV strain B (Figure 3). Amino acid alignments among the six NGPV strains (SDHT16, SDJN19, SDLC19, SDLC01, JS1, and QH15) and the classical GPV strains indicated that 12 amino acid mutations were produced in the NGPV strains compared with the classical GPV strains (Table 3). The altered amino acid sites were I50T, K131R, A140S, A350V, V468I, E498R, R553K, N555T, E573Q/K, D594Y, N609D, and V617A.

Sequence comparison of the VP1 proteins

The VP1 gene of three isolates was composed of 2199 nucleotides, coding for 732 amino acids. Homology comparison indicated that the VP1 proteins of three NGPV strains shared 99.1%–99.7% nucleotide similarities between each other but showed 94.4%–95.1% similarities with the Chinese GPV strains (82-0321, 06-0329, YZ99-6, and LH) and 96.0%–96.6% similarities with the European GPV strain B (Figure 3).

Amino acid alignments of the VP1 proteins among the six NGPV strains (SDHT16, SDJN19, SDLC19, SDLC01, JS1, and QH15) and the classical GPV strains indicated that compared with the Chinese GPV strains, the six NGPV strains harbored 16 common amino acid alterations, among which nine mutated amino acid sites were identical with the European strain B (Table 4). The 16 altered amino acid sites were H28Q, T35S, A41D, Q89L, D114H, Q116H/Y, D142E, V144I, A180V, L207M, A/T210S, S450N, A524G, L537I, K575R, and D703N.

Sequence comparison of ITRs

The ITRs of three NGPV strains (SDHT16, SDJN19, and SDLC19) shared 98.1%–100.0% nucleotide sequence homologies but displayed 93.0%–94.0% homologies with the classical GPV strain LH. The alignments of ITR sequences revealed that strain SDHT16, SDJN19, and SDLC19 have additional two, four, and four nucleotide insertions in the stems of the palindrome region, respectively, compared with the classical GPV strain LH (Fig. 4).

Phylogenetic tree construction

Because only partial sequences of the European SBDS-related NGPV strains were sequenced, a 427-bp fragment at the N terminus of the VP1 gene was used to construct the phylogenetic tree. The results revealed that six Chinese SBDS-related NGPV, including SDHT16, SDJN19, SDLC19, SDLC01, JS1, and QH15, intimately clustered together and grouped with the European SBDS-related NGPV strains (D697/3/06 and D146/02), forming a separate branch (Fig. 5). The four Chinese GPV strains (LH, YZ99-6, 82-0321, and 06-0329) closely clustered together and grouped with two European GPV strains (VG32/1 and B), forming another separate branch. On the basis of the results of phylogenetic analysis, the

Chinese SBDS-related NGPV probably evolved from a common ancestor with the European SBDS-related NGPV.

Discussion

Viral isolation and PCR characterization are the preferred methods of diagnosing SBDS cases. In contrast to the parvoviral disease of geese (Derzsy's disease), against which the GPV isolation is easily achieved when using embryonated goose eggs as the vector [5, 6, 15], viral isolations against SBDS cases are not always successful [1, 14, 20]. The primarily isolated virus may be lost in the following blind passages in duck embryos; hence, the isolation rates of positive samples were comparatively low. In this study, goose embryos were used for viral isolation against SBDS cases. Although only a limited number of embryonated goose eggs died post-inoculation of positive samples, the primary isolates can stably kill the embryonated goose eggs in a shorter time in the next passaging stages. The results demonstrated that embryonated goose eggs still are a suitable platform for isolating NGPV from SBDS cases. However, it should be noted that classical GPV usually kills 12-day-old goose embryos between 3 and 7 days post-inoculation of tissue sample from Derzsy's disease [6], implying that classical GPV possesses a better replication capability in goose embryos than NGPV.

Strain JS1, QH15, and SDLC01 were previously isolated by other teams in China in 2015 [1, 12, 20]. The isolation year of SDHT16 was closer to that of strain JS1, QH15, and SDLC01 than SDJN19 and SDLC19. The VP1 and Rep1 proteins of SDHT16 displayed 99.8%–99.9% nucleotide sequence identities with that of strain JS1, QH15, and SDLC01, but showed 99.1%–99.4% and 99.3%–99.5% nucleotide sequence identities with that of strain SDJN19 and SDLC19, respectively. The result demonstrated that strain SDHT16 has the closer relationship with the early NGPV isolates than the 2019 NGPV isolates.

Strain SDJN19 and SDLC19 shared the higher genomic sequence homology (99.8%) than with SDHT16 (99.1% and 99.0%, respectively). In order to evaluate whether the 2019 NGPV isolates may have different replication ability from SDHT16, further comparisons were conducted between strain SDJN19 and SDHT16 in this study. The result demonstrated both strains can be passaged in embryonated cherry valley duck eggs, however, on the basis of the ELD₅₀ and MDT values comparisons, strain SDJN19 was found to possess a better propagation ability in embryonated cherry valley duck eggs than SDHT16. Given that the classical GPV cannot propagate in embryonated cherry valley duck eggs, we deduce that the minor nucleotide differences across the genome probably contribute to the differential propagation capability observed in strain SDJN19 and SDHT16. Whether strain SDJN19 and SDHT16 can exhibit the differential pathogenicity in cherry valley duck deserves to be investigated.

ITR acts as the replication origin of the genome, containing a number of transcription factor binding sites and short nucleotide repeats (TCCGGT) [19, 21]. Compared with the classical GPV strain LH, nucleotide mutations were produced in the ITR region of three NGPV isolates. ITR sequences of two 2019 NGPV isolates (SDJN19 and SDLC19) shared 99.5% similarities, but they showed a slightly low similarities (97.6%–98.1%) with SDHT16. Besides, three NGPV isolates harbored one or two additional nucleotide

insertions in the stem region of ITR. However, all of these nucleotide mutations and insertions were found to be located in the base-pairing positions, hence the correct hairpin structure formation of ITR was not influenced. Altogether, the sequence comparison results of ITRs among three NGPV strains demonstrated a closer relationship between two 2019 NGPV isolates, which was in parallel with the comparison results of the coding proteins.

The characteristic amino acids sites belonging to the NGPV isolates were identified in this study. These site alterations produced in the NGPV strains played a key role in the host transition from goose to cherry valley duck. According to our observation, the amino acid alterations of both proteins were not evenly distributed in the peptide sequences. In terms of Rep1 protein, 8 of 12 site alterations were located within one third of the peptide sequences adjacent to the carboxyl terminus. C termini of the Rep1 protein contains the domain involved in transactivation of P41 promoter, which modulate the VP1 gene transcription [13]. For the VP1 protein, half of 16 alterations were located in the VP1u region of the VP1 protein. VP1u contains the phospholipase A2 (PLA2) domain, which is required for parvovirus entry and infectivity [8, 22]. Further analysis showed only an amino acid alteration (Q89L) was found to be located in the PLA2 motif (53-111 aa) of VP1u of GPV [22], other seven mutated sites in the VP1u lay outside of the PLA2 motif. Nevertheless, the amino acid residues outside the PLA2 motif are also important for maintaining a proper three dimensional structure that is essential for the PLA2 activity [4]. Together, the common amino acid mutations identified in this study should contribute to the host range shift of NGPV.

Sequence alignment and comparison revealed that the six Chinese NGPV isolates have closer relationship with the European strain B than with the Chinese GPV strains. In the phylogenetic tree, the Chinese NGPV isolates grouped with the European SBDS-related NGPV strains and formed a separate branch, distinct from the branch grouped by the classical GPV strains. The present analysis result indicates that the Chinese NGPV isolates probably originate from a common ancestor with the European counterpart. The European SBDS-related NGPV was isolated from diseased mule ducks, which are a sterile and intergenetic cross of Pekin and Muscovy ducks [14]. Therefore, whether a characteristic difference exists at the genome level between the European and Chinese SBDS-related NGPV isolates needs to be elucidated. Whole genome sequencing against the European SBDS-related NGPV isolates may be able to shed light on this matter.

Taken together, three new SBDS-related NGPV isolates were obtained in this study, and their genomes were sequenced. The differential propagation abilities in embryonated cherry valley duck eggs were observed between two NGPV strains isolated in different years. The characteristic amino acid differences between the NGPV strains and the classical GPV strains were identified, which should contribute to host range shift. Further phylogenetic analysis supported the conclusion that the Chinese NGPV isolates probably originate from a common ancestor with the European SBDS-related NGPV.

Last but not least, the VP1 protein of NGPV still shares almost 95% amino acid identity with the classical GPV, irrespective of the characteristic amino acid differences. The structural proteins, including VP1, VP2, VP3, commonly constitute the protective antigens of GPV [7]. Hence, whether the traditional GPV vaccine

currently in use against the Derzsy' disease can be applied in cherry valley ducks to prevent from SBDS deserves to be thoroughly investigated.

Declarations

Funding

This work was supported by the National Natural Science Foundation of China [grant numbers 31572551, 31172317] and by a project funded by the Priority Academic Program Development of Jiangsu Higher Education Institutions.

Conflicts of interest/Competing interests

The authors declare that they have no competing interests.

Ethics approval

The procedure for inoculation of embryonated goose or duck eggs was approved by the Animal Care and Use Committee of Yangzhou University and performed in accordance with the "Guidelines for Experimental Animals" of the Ministry of Science and Technology (Beijing, China). No specific permissions were required for these locations/activities.

Consent to participate (include appropriate statements)

All the listed authors participated in this study with informed consents

Consent for publication (include appropriate statements)

Written informed consent for publication was obtained from all participants.

Availability of data and material

Not applicable

Code availability

Not applicable

Authors' contributions

Li Y, Jia J, and Mi Q carried out the overall experiments and analyzed the data. Li Y, and Gao Y collected the tissue samples during the SBDS disease outbreak in cherry valley duck farms. Li Y, Jia J wrote the paper. Zhu G helped to analyze the result and contributed to language polishing. Wang J conceived the project and directed the research.

Acknowledgements

This work was supported by the National Natural Science Foundation of China [grant numbers 31572551, 31172317] and by a project funded by the Priority Academic Program Development of Jiangsu Higher Education Institutions.

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Tables

Table 1 Primers designed for PCR amplification of the whole genomes of the NGPV isolates

Primer numbers	Primer sequences (5'→3')	Primer positions	Fragment sizes (bp)
ITR-1	tcattggagggttcgttcg	1–23; 5024–5046	188
ITR-2	gcatgCGCGCGGTCaacctaa	168–188; 4859–4879	
ITR-3	gcatgCGCGGTCagcccaa	183–203	476
ITR-4	ctccaaactccggtagaacc	622–644	
P1	atggcactttctaggcctct	507–526	1378
P2	aagaaggctccaatttatgtga	1863–1884	
P3	gtattgaacctactcctgtaatt	1741–1763	1428
P4	tgtagatgtggttgtgtagc	3148–3168	
P5	ccgatggagtgggtaatgcct	3049–3069	1550
P6	ttgagtagatatctggtccaat	4575–4598	
ITR-5	ggccagtgtacagtagagatggtg	4425–4448	440
ITR-3	gcatgCGCGGTCagcccaa	4844–4864	

Table 2 Background of the classical and novel goose parvovirus strains used for sequence comparison and phylogenetic analysis

Strain	Host	Geographic origin	Isolation year	Pathogenicity	GenBank accession number
B	Goose	Hungary	1967	Pathogenic	U25749
VG32/1	Goose	Germany	n.a.	Vaccine	EU583392
YZ99-6	Goose	China	1999	Pathogenic	KC996730
LH	Goose	China	2003	Pathogenic	KM272560
82-0321	Goose	China	1982	Pathogenic	EU583390
06-0329	Goose	China	2006	Pathogenic	EU583391
SDLC01	Cherry valley duck	China	2015	Low pathogenic	KT343253
JS1	Cherry valley duck	China	2016	Low pathogenic	KT935536
QH15	Cherry valley duck	China	2015	Low pathogenic	KT751090
SDHT16	Cherry valley duck	China	2016	Low pathogenic	MN356043
SDJN19	Cherry valley duck	China	2019	Low pathogenic	MN356044
SDLC19	Cherry valley duck	China	2019	Low pathogenic	MN356045
D146/02	Mule duck	France	2002	Low pathogenic	AY496906
D697/3/06	Mule duck	France	2006	Low pathogenic	EU938706

n.a.: not available

Due to technical limitations, table 3,4 is only available as a download in the Supplemental Files section.

Figures

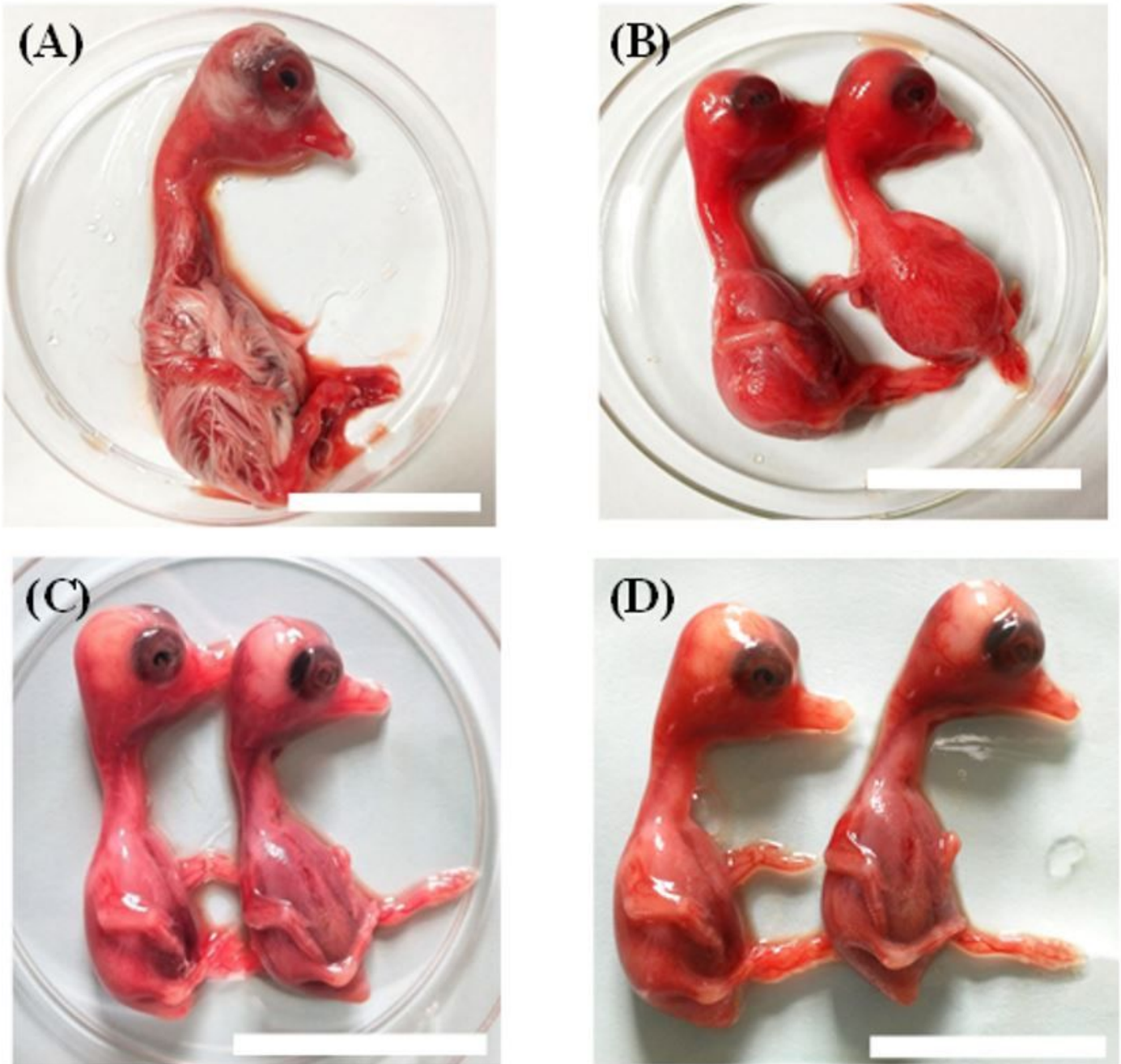


Figure 1

Hemorrhagic lesions were observed in goose or duck embryos that died at different time points post-infection during the isolation and passage stage of strain SDJN19 and SDHT16. (A) Goose embryos died at 264 h post-inoculation of the tissue sample homogenate. (B) Goose embryos died at 112 h after the first passage with the SDJN19 isolate. (C) Cherry valley duck embryos died at 96 h post-infection of the SDJN19 isolate. (D) Cherry valley duck embryos died at 112 h post-infection of the SDHT16 isolate. The scale bar is 30 mm.

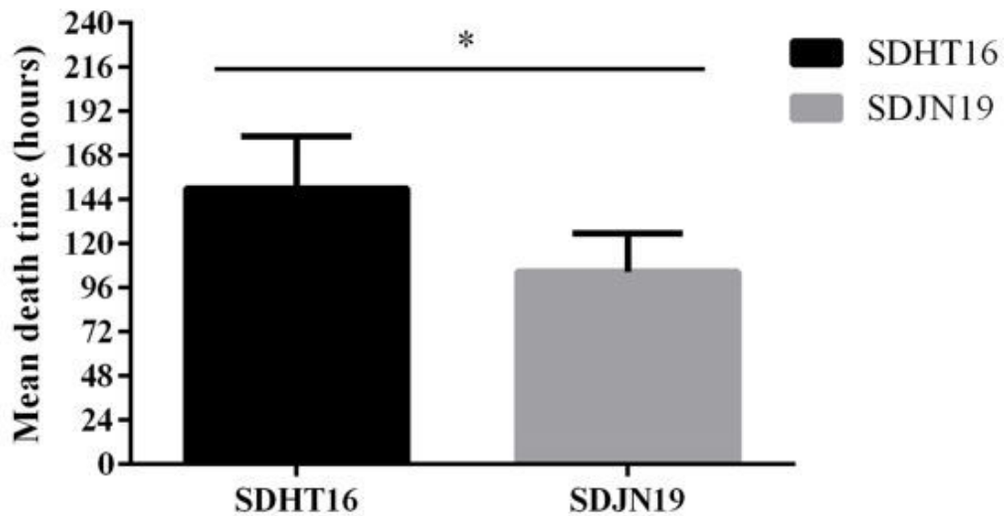


Figure 2

Comparison of the mean death time of the embryonated cherry valley duck eggs post-infection of strain SDHT16 and SDJN19.* Significant difference between SDHT16 and SDJN19-infected groups (P < 0.05).

		Percent identity of VP1												
		1	2	3	4	5	6	7	8	9	10	11		
Percent identity of Rep1	1		96.5	96.7	96.4	96.6	96.7	96.6	96.4	96.3	96.0	96.6	1	B
	2	99.3		99.6	99.3	99.4	95.0	94.9	94.7	94.7	94.5	94.9	2	LH
	3	99.2	99.8		99.4	99.6	95.2	95.1	94.9	94.9	94.7	95.1	3	YZ99-6
	4	98.9	99.4	99.3		99.6	94.9	94.8	94.6	94.6	94.4	94.8	4	06-0329
	5	99.4	99.8	99.7	99.4		95.1	95.0	94.8	94.7	94.5	95.0	5	82-0321
	6	96.9	96.4	96.3	96.1	96.4		99.9	99.7	99.3	99.0	99.9	6	SDLC01
	7	96.9	96.4	96.3	96.1	96.5	99.8		99.7	99.4	99.1	99.9	7	JS1
	8	96.9	96.4	96.3	96.1	96.5	99.8	100.0		99.2	98.9	99.8	8	QH15
	9	96.9	96.4	96.3	96.1	96.5	99.4	99.6	99.6		99.7	99.4	9	SDJN19
	10	96.8	96.3	96.2	96.0	96.4	99.2	99.3	99.3	99.7		99.1	10	SDLC19
	11	96.9	96.4	96.3	96.1	96.4	99.8	99.9	99.9	99.5	99.3		11	SDHT16
		1	2	3	4	5	6	7	8	9	10	11		

} SBDS-related NGPV

Figure 3

Percentages of the identities shared by the nucleotide sequences of the coding proteins VP1 (upper right) and Rep1 (lower left) of six NGPV strains and five classical GPV strains. Percentages of identities among the six NGPV strains are shaded in gray.

LH	TCATTGGAGGGTTCGTTTCGTTCCGAACCCAGCCAATCAGGGGAGGGGG- ⁴⁷ AAGTGACGCAAGTTCGGGTGACGCACATCCGGT	79
SDHT16	TTATTGGAGGGTTCGTTTCGTTCCGAGCCAGCCAATCAGGGGAGGGGG-AAGTGACGCAAGTCCGGTCAAGTGCCTTCGGT	79
SDJN19	TTATTGGAGGGTTCGTTTCATTCCGAGCCAGCCAATCAGGGGAGGGGGGAAGTGACGCAAGTTCGGGTGACGTGCCTTCGGT	80
SDLC19	TTATTGGAGGGTTCGTTTCATTCCGAGCCAGCCAATCAGGGGAGGGGGGAAGTGACGCAAGTTCGGGTGACGTGCCTTCGGT	80
113		
LH	GACGTAGTTCGGGTACAGTGCCTTCCTGTACAGTG-TTCCGGTACAGTGACTTCGGTTCATGTGACTTCCGGTGACGTGT	158
SDHT16	GACGCACATCCGGTACAGTGCCTTCCTGTACAGTGACTTCGGTTCATGTGACTTCGGTTCATGTGACTTCCGGTGACGTGT	159
SDJN19	GACGCACATCCGGTACAGTGCCTTCCTGTACAGTGACTTCGGTTCATGTGACTTCGGTTCATGTGACTTCCGGTGACGTGT	160
SDLC19	GACGCACATCCGGTACAGTGCCTTCCTGTACAGTGACTTCGGTTCATGTGACTTCGGTTCATGTGACTTCCGGTGACGTGT	160
Bubble region		
LH	TTCCGGCTGTTAGGTTGACCGCGCGCATGCGCGCGGTCAGCCCAATAGTTAAGCCGGAAACACGTACCCGGAAGTCACAT	238
SDHT16	TTCCGGCTGTTAGGTTGACCGCGCGCATGCGCGCGGTCAGCCCAATAGTTAAGCCGGAAACACGTACCCGGAAGTCACAT	239
SDJN19	TTCCGGCTGTTAGGTTGACCGCGCGCATGCGCGCGGTCAGCCCAATAGTTAAGCCGGAAACACGTACCCGGAAGTCACAT	240
SDLC19	TTCCGGCTGTTAGGTTGACCGCGCGCATGCGCGCGGTCAGCCCAATAGTTAAGCCGGAAACACGTACCCGGAAGTCACAT	240
262		
LH	GACCCGGAAGTCACGTGACCCGAAA-CACGTGACAGGAAGCACGTGACCCGAACTACGTACCCGGATGTGCGTCACCCGAA	317
SDHT16	GACCCGGAAGTCACATGACCCGAAAGTACAGTACAGGAAGCACGTGACCCGAACTACGTACCCGGATGTGCGTCACCCGAA	319
SDJN19	GACCCGGAAGTCACATGACCCGAAAGTACAGTACAGGAAGCACGTGACCCGAACTACGTACCCGGATGTGCGTCACCCGAA	320
SDLC19	GACCCGGAAGTCACATGACCCGAAAGTACAGTACAGGAAGCACGTGACCCGAACTACGTACCCGGATGTGCGTCACCCGAA	320
334		
LH	CTTGCGTCACTTCCCCC-TCCCCTGATTGGCTGGTTCGAACGAACGAACCCTCCAAT ³⁷⁶ GAGACTCAAGGACAAGAGGATAT	396
SDHT16	CTTGCGTCACTTCCCCC-TCCCCTGATTGGCTGGTTCGAACGAACGAACCCTCCAATAAGACTCAAGGACAAGAAAATAT	398
SDJN19	CTTGCGTCACTTCCCCCCTCCCCTGATTGGCTGGTTCGAATGAACGAACCCTCCAATAAGACTCAAGGACAAGAAAATAT	400
SDLC19	CTTGCGTCACTTCCCCCCTCCCCTGATTGGCTGGTTCGAATGAACGAACCCTCCAATAAGACTCAAGGACAAGAAAATAT	400
D sequence		
LH	<u>TTTGCGCGCCAGGAAGTG</u>	414
SDHT16	<u>TTTGCGCGCCAGGAAGTG</u>	416
SDJN19	<u>TTTGCGCGCCAGGAAGTG</u>	418
SDLC19	<u>TTTGCGCGCCAGGAAGTG</u>	418

Figure 4

Nucleotide sequence alignment of ITRs among three NGPV strains (SDHT16, SDJN19, and SDLC19) and the classical GPV strain LH. The different nucleotides or insertions relative to the classical GPV strain LH are shaded in grey. The dashes denote deletions. The sequences in the bubble region of ITR are composed of 43 nucleotides and indicated by open circles above the letters. The D sequences are underlined, which reversely complement with the D' sequences at the 3' ITR.

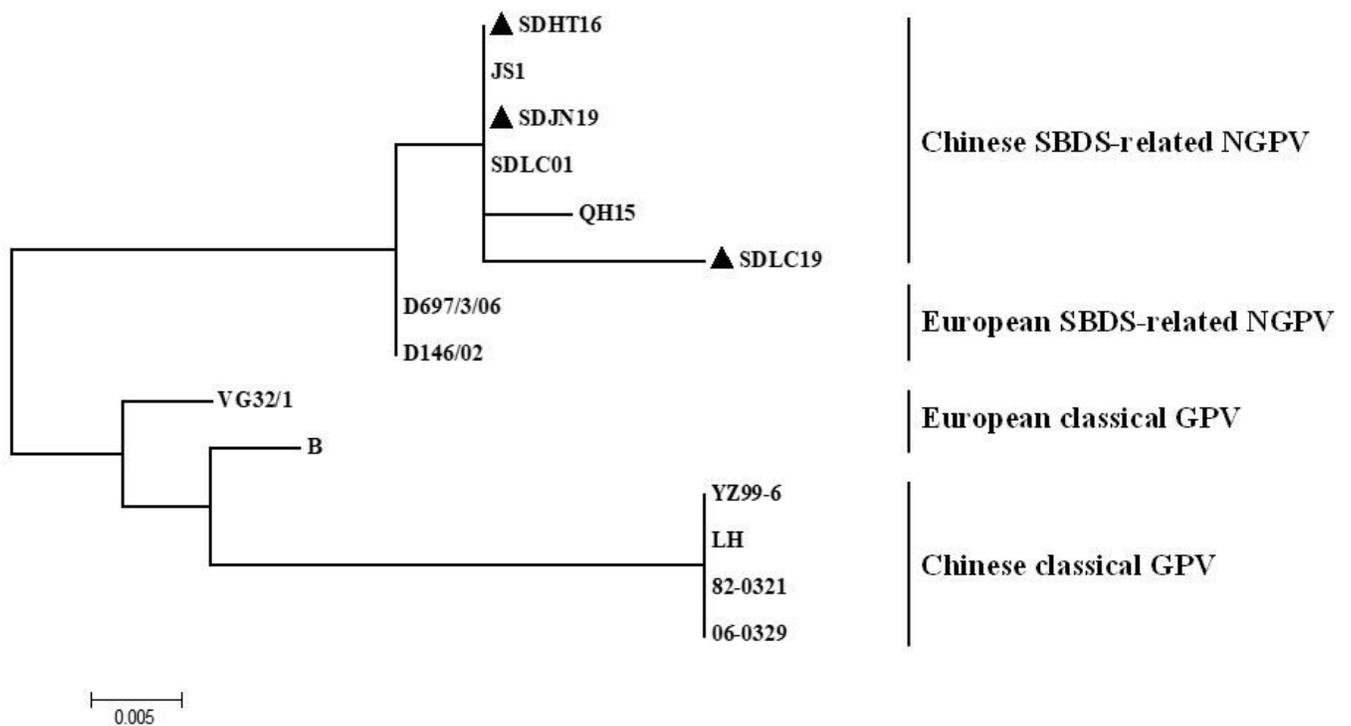


Figure 5

Phylogenetic analysis of the SBDS-related NGPV and the classical GPV strains. The phylogenetic tree was constructed on the basis of a 427-bp VP1 gene fragment. The Chinese SBDS-related NGPV strains clustered with the European NGPV strains, forming a separate branch. Phylogenetic trees were constructed through the neighbor-joining method in MEGA 7.0 with 1000 bootstrap replications.

Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.

- [Table3.jpg](#)
- [Table4.jpg](#)