Molecular Characterization of a Porcine Sapelovirus Strain Isolated in China

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Abstract

Background

Porcine sapelovirus (PSV) infections have been associated with a wide spectrum of symptoms ranging from asymptomatic infection to clinical signs including diarrhoea, pneumonia, reproductive disorders, and polioencephalomyelitis. Although it has a global distribution, studies on PSV in domestic have been relatively few so far which goes against for this pathogen's research and disease prevention.

Methods

In this study, we isolated a PSV strain, SHCM2019, from the swine faecal specimens with PK-15 cells. To investigate its molecular characteristics and pathogenicity, the genomic sequence of the SHCM2019 strain was analysed, and further the clinical manifestations and pathological changes exhibited following inoculation of the neonatal piglets were observed.

Results

The strain isolated with PK-15 cells, was identified as PSV by RT-PCR, IFA and TEM assays. Sequencing results showed that the full-length genome of the SHCM2019 strain was 7567 nucleotides including a 27-nucleotide poly (A) tail. Phylogenetic analysis demonstrated that the virus isolate belonged to PSV and was classified into the Chinese strain cluster. Virus recombination analysis indicated that there might be a recombination break point upstream of the 3D region in PSVs. Pathogenicity research demonstrated that the virus isolate could cause diarrhoea and pneumonia in piglets.

Conclusion

This study presented the isolation of a recombinant PSV strain, SHCM2019, and demonstrated the isolate was pathogenic. Our results may provide a reference for future researches of the pathogenic mechanism and evolutionary characteristics of PSV.

Introduction

The genus Sapelovirus, belonging to the family Picornaviridae, consists of three species: Avian Sapelovirus, Sapelovirus A (porcine sapelovirus), and Sapelovirus B (simian sapelovirus) [1]. Porcine sapelovirus (PSV) is a non-enveloped, positive-sense, single-stranded RNA virus, which was previously known as porcine enterovirus 8 (PEV-8) in the genus Enterovirus [2]. The genome of PSV measures approximately 7500 base pairs (bp) in length, consisting of 5' untranslated region (UTR), intermediate coding region, 3' untranslated region (UTR) and poly (A) tail. The intermediate coding region contains only a large open reading frame (ORF) that encodes a polyprotein precursor about 2,331 amino acids in length, which is subsequently cleaved by the self-coding protease into 12 mature functional proteins. The structure of the polyprotein is consistent with the L-4-3-4 structure of other picornaviruses, in which a
leading peptide L leads to four structural polypeptides (VP4, VP2, VP3, and VP1) and seven nonstructural polypeptides (2A, 2B, 2C, 3A, 3B, 3C, and 3D) [3–5].

Since PSV was first reported in the UK in 1958 [6], it has been identified in many countries, including Canada, Japan, Australia, Brazil, Spain and Korea, and China and other countries worldwide [5, 7–12]. PSV infection rates in pigs have been investigated globally, ranging from 7.1% in India to 71.0% in Hungary [13, 14]. PSV infections have been associated with a wide spectrum of symptoms ranging from asymptomatic infection to acute fatal encephalomyelitis, reproductive disorders, diarrhoea and pneumonia [12, 15–17]. Prodelalova et al reported that PSV infection rates were higher in asymptomatic pigs than in diarrhoeal pigs [15]. However, Zhang et al demonstrated there was a higher prevalence of PSV in diarrhoeal pigs by using metagenomic analysis of pig faeces [18]. Further, Kim et al reported that PSV could cause intestinal lesions and colonize the villous epithelial cells in the small intestine [19]. Like porcine epidemic diarrhea virus (PEDV), PSV can coinfect with porcine parvovirus, swine fever virus, porcine reproductive and respiratory disorder syndrome virus, porcine enterovirus and other viruses, leading to clinical atypical symptoms in sick pigs [20–22]. the symptoms of PSV are often covered up and overlooked due to the mixed infections [15], which poses a considerable threat to the pig industry.

In this study, a PSV strain was isolated using PK-15 cells. Based on the phylogenetic analysis, it is closely related to other Chinese isolates. The recombination analysis indicated a possible recombination break point in the 3D region. Animal regression experiments demonstrated that the isolated strain was pathogenic to piglets. The study laid a foundation on the pathogenic mechanism of PSV.

**Materials And Methods**

**Cells and clinical samples**

PK-15 cells were cultured at 37°C in 5% CO\(_2\) in Dulbecco's modified Eagle's medium (DMEM, HyClone, USA) containing 10% heat-inactivated foetal bovine serum (FBS; Invitrogen, Australia) and 1% antibiotics (10,000 units/mL penicillin and 10,000 µg/mL streptomycin). The faecal specimens were collected from the diarrheic piglets from a pig farm in Shanghai, China, in 2019. Subsequently, these faecal samples were used for multiplex PCR detection of several potential pathogens including PSV, PEDV, porcine kobuvirus (PKV), porcine astrovirus (PAstV), porcine deltacoronavirus (PDCoV), porcine torovirus (PToV), porcine teschovirus (PTV), porcine sapovirus (PSaV), transmissible gastroenteritis virus (TGEV), porcine rotavirus (PoRV) and bovine viral diarrhoea virus (BVDV).

**Virus isolation**

PK-15 cells were inoculated with faecal specimens that were RNA-positive for PSV but RNA-negative for PEDV, TGEV, PKV, PAstV, PDCoV, PToV, PTV, PSaV, PoRV, and BVDV. In brief, the positive faecal samples were suspended in sterile phosphate-buffered saline (PBS) containing 1,000 units/mL penicillin and 1,000 µg/mL streptomycin and centrifuged for 10 min at 3,500×g. The supernatants were filtered through 0.22-µm pore membrane filters (Merck Millipore Ltd., USA), followed by inoculation in PK-15 cells for 1 h.
at 37°C. Next, the inoculum was discarded, and the cells were washed three times with sterile PBS. Subsequently, DMEM supplemented with 0.5 µg/mL TPCK-trypsin (Gibco™, USA) was added, and cell culture was continued at 37°C in 5% CO₂, with daily observation for cytopathic effect (CPE). When more than 80% of cells appeared CPE, the supernatant and cells were harvested for continuous subculture.

**RT-PCR**

Total RNA of the first six generations was extracted from the culture supernatant of PSV inoculation using TRIzol Reagent (Takara, China) according to the manufacturer’s instructions. The cDNAs were synthesized in a final volume of 20 µL containing 10 µL RNA, 4 µL 5×RT buffer, 1 µL dNTPs (10 mM), 2 µL random primer (25 µM), 0.5 µL Moloney murine leukaemia virus (M-MLV) reverse transcriptase (TaKaRa, China), 0.5 µL RNase inhibitor (TaKaRa, China) and 2 µL diethylpyrocarbonate (DEPC)-treated water. The reaction was incubated at 42°C for 1h followed by incubation at 70°C for 15 min. cDNA was used for PCR with the following primers, PSV-F:5′-GATGTGGCGCATGCTCTT-3′, PSV-R:5′-TGCTGCTCCTGTTGTTTTAT-3′. Amplified products were separated by electrophoresis in a 1.5% agarose gel and purified using a DNA gel extraction kit (Tiangen, China). The purified PCR products were subsequently cloned into the PEASY-Blunt Zero vector (TransGen, China) and sequenced (Biosune, China). Sequences were analysed using the BLAST search program (http://blast.ncbi.nlm.nih.gov/).

**Transmission electron microscopy (TEM)**

The PK-15 cells infected with strain SHCM2019 were frozen and thawed thrice when there were over 80% CPE, followed by centrifugation at 2,000×g for 30 min to remove cell debris. Crude virus was pelleted from the clarified supernatant by ultracentrifugation at 9,890×g at 4°C for 3 h in a TYPE-70 Ti rotor (Beckman, USA). The resulting pellet was resuspended in a small amount of PBS and was subsequently layered onto a 20–60% (w/v) discontinuous iodixanol solution (OptiPrep™ Density GradientMedium, Sigma-Aldrich, USA) by centrifugation at 124,400×g at 4°C for 3 h in a SW-55 Ti rotor (Beckman, USA). The virus band at the interface was collected and placed onto a formvar grid (Electron Microscopy Sciences, USA) for 5 min, and excess liquid was subsequently removed by filter paper. The samples were observed using a transmission electron microscope (Hitachi, Japan) for the morphological identification of the virus.

**Immunofluorescence assay (IFA)**

To characterize the isolated strain, IFA was performed with a polyclonal antibody specific for the PSV-VP1 protein (prepared in our laboratory). Briefly, PK-15 cells in 6-well plates were mock infected or infected with the SHCM2019 strain, incubated for 24 h, fixed in 4% paraformaldehyde for 30 min at 4°C, and subsequently permeabilized with 0.3% Triton X-100 (Biodee, China) for 10 min at room temperature (RT). After the cells were washed three times with PBS, they were blocked with 5% bovine serum albumin (BSA; Sigma, USA) at RT for 1 h. Mouse polyclonal antibody against PSV and Alexa Fluor™ 488-conjugated rabbit anti-mouse IgG (Invitrogen, USA) were used as primary and secondary antibodies, respectively. The cell nuclei were stained with 4′,6-diamidino-2-phenylindole (DAPI; Beyotime, China) for 5 min at RT. After
the cells were washed with PBS, the stained cells were observed with a fluorescence microscope (Carl Zeiss, Germany).

**Sequence and phylogenetic analysis**

Third-generation viral RNA was extracted from PK-15 cells infected with the virus isolate using TRIzol Reagent. Reverse transcription and PCR were performed with M-MLV reverse transcriptase and TransStart FastPfu DNA polymerase (TransGen, China), respectively. The PCR products were cloned using the pEASY-Blunt Cloning Kit, and the resulting plasmids were sequenced. The 5'-UTR and 3'-UTR sequences of the isolated strain were obtained using a 5'-3' RACE kit (Takara, China) according to the manufacturer's instructions. The sequences fragments were assembled with DNAStar v. 7.1. Subsequently, the complete sequence was submitted to GenBank (GenBank ID: MN685785). All of the complete sequences of the polyprotein gene and reference sequences obtained from GenBank were utilized in sequence alignments and phylogenetic analyses. Phylogenetic trees were constructed using the neighbour-joining method with 1000 bootstrap replications in MEGA v. 7.0. Recombination analysis was performed using SimPlot v. 3.5.1 and Recombination Detection Program (RDP) v. 4.10.

**Experimental infection of piglets**

To assess the pathogenicity of the isolated SHCM2019 strain in piglets, four 5-day-old piglets from the same sow were used. Before inoculation, we observed for 3 days and collected faecal samples from all piglets to ensure that none of the piglets were infected with PSV. The four piglets were random divided into two groups (two piglets per-group). Each piglet in group 1 (L1 and L2) was given 5 mL PSV-SHCM (10^{7.41} TCID_{50}/mL) by the oral route, and piglets in group 2 (C1 and C2) received 5 mL DMEM orally as a negative control. After inoculation, all piglets were monitored each day for clinical signs. Faecal samples of each piglet were collected for viral load detection by qRT-PCR at 0, 1, 2, 3, 4, 5 and 6 days post-inoculation (dpi), and serum samples were collected for seroconversion tests at 0, 1, 3 and 6 dpi. All piglets were artificial executed at 6 dpi, and one portion of the tissue samples were fixed with 4% paraformaldehyde solution for histological slides preparation and the other portion were collected for viral load detection by qRT-PCR.

**Ethics statement**

All animal experiments were performed under the guidance of the Institutional Animal Care and Use Committee at the Center for Disease Control and Prevention (CDC) and the Laboratory Animal Care International accredited facility.

**Results**

**Virus isolation and identification**

A PSV strain was isolated from diarrheal faecal samples originating from a farm in Shanghai, China. PK-15 cells infected with the isolate displayed a distinct CPE after three times of passages, which was
characterized by rounding and shrinking of the cells after 30 h and all cells fell off at 70 h (Fig. 1A). RT-PCR assays with a pair of primers specific to PSV amplified a 624 bp target fragment (Fig. 1B). Further IFA confirmed that the polyclonal antibody against PSV could be reacted with the SHCM2019 strain with the specific green fluorescences in cytoplasm (Fig. 1C). To determine its morphological characteristics, PSV particles of the SHCM2019 strain were purified and stained negatively, following imaged by transmission electronmicroscopy (TEM). It was revealed that the viral particles were nonenveloped, icosahedral, approximately 25–30 nm in diameter (Fig. 1D).

Full genome organization and phylogenetic analyses

To further explore the characteristics of the isolated virus, the complete genomic sequence of strain SHCM2019 was tested. The full-length genome of this virus strain was 7,567 nucleotides (nt) with only a single 6996-nt ORF flanked by a 465 nt 5′UTR and a 106 nt 3′ UTR. Although the genome of strain SHCM2019 in length is different from other PSV strains, its polyprotein gene is similar to most Chinese PSVs. Actually, the length of the polyprotein gene is varied in PSVs, however, it is relatively conserved in Chinese PSVs, exhibiting a consistent length of 6996 nt, except for the HuN4, HuN6, HuN21, and HuN32 strain with some extra nucleotides inserted in it.

The phylogenetic tree was constructed by the neighbour-joining method based on the complete polyprotein gene of the SHCM2019 strain and other representative picornaviruses from the NCBI database. The results indicated that SHCM2019 was most similar with the Chinese PSV strains, which were clustered in the same group (Fig. 2). Large genetic diversity was existed among the PSVs isolated from different countries. The Chinese PSV strains were closely related to the Korean PSV strains. In addition, the SHCM2019 strain showed high nucleotide (85.8–91.2%) and amino acid (71.5–79.8%) identities with the other PSV strains, but relatively low identities with the avian and simian sapelovirus strains (Supplemental Fig. 1).

Further the virus recombination analysis was carried out using SimPlot v. 3.5.1 and RDP v. 4.10 softwares. The SHCM2019 was used as separate queries to implement a standard similarity plot analysis. Next, we performed a bootscanning analysis to verify potential breakpoints in the recombinants. The SHCM2019 strain showed a high degree of nucleotide sequence similarity with HeB04 in the VP1, VP2 and VP3 regions, whereas the 3D regions were highly similar to those of ISU-SHIC, which suggested that there might also be a restructuring break point in 3D (Fig. 3A, B).

Experimental infection, necropsy, and histopathological examination

To test the virulence of the SHCM2019 strain, two 8-day-old healthy piglets were inoculated with SHCM2019. Inappetence and lethargy appeared in PSV-infected piglets at 2 dpi, and watery diarrhoea occurred at 5 dpi compared to the mock piglets (Fig. 4D). Necropsy results showed that lesions were mainly concentrated in the intestines and lungs. The intestines of PSV-infected piglets were thin and
translucent, and the intestinal contents were pulpy or watery (Fig. 4E). There was obvious congestion in the lungs compared to uninfected pigs (Fig. 4F). The piglets in the control group were clean around anus and had no obvious clinical symptoms (Fig. 4A). The intestines were thick and elastic, and the lungs were normal (Fig. 4B,C). Diseased tissues were also examined histopathologically. Histopathological results showed that villi in the small intestine were slightly atrophic, and submucosal oedema was obvious (Fig. 5C). The lung was damaged by thickened alveolar walls, serous exudation and inflammatory cell infiltration in the interalveolar septa (Fig. 5D). There was no significant change in the control group (Fig. 5A,B).

To explore the excretion and distribution of PSV, viral load detection of faecal samples collected daily and tissue samples obtained after necropsy of piglets were conducted by qRT-PCR. The results showed that there were high RNA copy numbers in faecal samples at 3–6 dpi (Fig. 6A). In addition, PSV was determined to be widely distributed in lung and intestinal tissues. However, PSV wasn’t detected in the heart, liver, or kidney (Fig. 6C, D). The results of enzyme-linked immunosorbent assay (ELISA) showed that the IgG antibody levels of PSV-infected pigs were significantly higher than those of uninfected pigs, and antibody levels increased significantly at 6 dpi (Fig. 6B). These results confirmed the pathogenicity of SHCM2019 to piglets.

Discussion

PSV is widely popular among pig population in China. In this study, a PSV isolate from pig farms in Shanghai, SHCM2019, was identified and sequenced. The isolated strain has 7,567 nucleotides, consisting of a 5’ UTR of 465 nucleotides, a 3’ UTR of 106 nucleotides, and a single ORF of 6,996 nucleotides that encodes a polyprotein precursor of 2,331 amino acids. The length of the 5’ UTR is varied in PSVs. According to previous research, the 5′ terminal residues of the complete picornavirus should be UU [5], but the two 5′ terminal residues were AC for strain SHCM2019, which was not accordance with those sequence features. The antigenic epitome of VP1 had high identity with other PSV strains, indicating the slow evolution, which will provide useful clues for disease prevention.

Phylogenetic analysis based on the polyprotein gene revealed that the PSV isolates from China exhibit genetic diversity, and are closely related to the South Korean strains, but are distantly related to the PSV isolates from Europe and America, suggesting that the prevalence of PSVs may has certain territoriality. Furthermore, we constructed an evolutionary tree of the VP1 gene (Supplementary Fig. 2). However, the topological structure and phylogenetic branching of the VP1 tree were slightly different from the ORF tree and the VP1 tree previously reported [23], which indicated that PSV could not be divided into genotypes at present. At present, many picornaviruses are genotyped by VP1 sequence alignment[24]. The classification criterion is that the VP1 sequence difference between isolates is greater than 25% [25]. Among PSVs, the nucleotide and amino acid sequence identities of the VP1 region were > 70.8% and > 75.1%, respectively, and these values were less multiple than those of porcine enteroviruses (> 53.9% and > 53.3%, respectively) [24]. This might be one of the reasons why the genotyping PSVs is considered to be hard.
The driving forces of viral evolution include gene mutation, natural selection, gene drift, and viral gene recombination. In which, mutation and recombination are two important mechanisms. The results of recombination analysis showed recombination existed among the PSV isolates, however, the recombination site was located in the 3D region, which was not consistent with the previous studies [23, 26–28]. The 3D protein is a virus-specific RNA polymerase[3], which has the ability to synthesize viral RNA, and its recombination plays an important role in viral replication. In view of the diversity of recombination sites reported to date [23, 26–28], we must accelerate the process of research on the molecular mechanism of PSV.

The SHCM2019 strain was found to replicate in the intestinal tracts of piglets and cause diarrhoea and a degree of pneumonia in pigs, suggesting that the SHCM2019 strain is pathogenic. In clinic, like porcine circovirus type 2 (PCV2), PSV infection rate was high in both domestic and wild pigs [9], but in most cases, the pigs infected with PSV did not exhibit clinical symptoms [8, 9, 14, 20], which suggests that PSV is not consistently pathogenic. This also led to PCV2 and PEDV being the main focus of most researchers when large-scale diarrhea occurred in pig populations, while, PSV, which could cause similar symptoms, didn't attract similar attention [29, 30]. Perhaps PSV require currently unidentified factors to show full pathogenicity in pigs, which warrant further investigation.

**Conclusion**

In this study, a swine sapelovirus strain was isolated and named SHCM2019. The isolated strain was confirmed as PSV by RT-PCR, IFA and TEM assays. The molecular characteristics and pathogenicity of the SHCM2019 strain were studied. Phylogenetic analyses showed that the virus isolate belongs to PSV and was classified into the China cluster. Recombination analysis indicated that there may be a recombination breakpoint upstream of the 3D region in PSVs. Pathogenicity research demonstrated that the SHCM2019 strain is pathogenic. To better assess the adverse effects of PSV on pig populations, we need to further explore the epidemiological characteristics and pathogenic mechanisms of PSV.

**Abbreviations**

PSV: Porcine sapelovirus; PEV-8: Porcine enterovirus 8; UTR: Untranslated region; ORF: Open reading frame; DMEM: Dulbecco's modified Eagle's medium; FBS: Foetal bovine serum; PKV: Porcine kobuvirus; PAstV: Porcine astrovirus; PDCoV: Porcine deltacoronavirus; PToV: Porcine torovirus; PTV: Porcine teschovirus; PSaV: Porcine sapovirus; PEDV: Porcine epidemic diarrhoea virus; TGEV: Transmissible gastroenteritis virus; PoRV: Porcine rotavirus; BVDV: Bovine viral diarrhoea virus; PCV2: porcine circovirus type 2; CPE: Cytopathic effect; PBS: Phosphate-buffered saline; M-MLV: Moloney murine leukaemia virus; DEPC: Diethylpyrocarbonate; TEM: Transmission electron microscopy; IFA: Immunofluorescence assay; RT: Room temperature; BSA: Bovine serum albumin; DAPI: Diamidino-phenylindole; RDP: Recombination Detection Program; ELISA: Enzyme-linked immunosorbent assay.

**Declarations**
Acknowledgements

Not applicable.

Author Contributions

HLL conceived and designed this study; NNL and BQL conducted experiments; JT, JHC and YS contributed their technical knowledge; NNL and HLL wrote this paper; BQL and TJ were involved in interpreting the results and critically reading the manuscript. All authors read and approved the final manuscript.

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Availability of data and materials

Data supporting the conclusions of this article are presented in the manuscript.

Ethics approval and consent to participate

All animal experiments were performed under the guidance of the Institutional Animal Care and Use Committee at the Center for Disease Control and Prevention (CDC) and the Laboratory Animal Care International accredited facility.

Consent to publication

Not applicable.

Competing interests

All authors declared that they have no competing interests.

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Isolation and identification of PSV. (A) Cytopathic effects in PK-15 cells inoculated with PSV-positive samples. After three times of passages, PK-15 cells began to shrink and shed at 30 h, and were widely detached from the surface of the cultivation flask at 70 h. (B) RT-PCR detection of cell culture medium obtained through continuous passages. The first six passages of viruses yielded positive bands of ~624 bp in RT-PCR analysis with primers specific to PSV. (C) Immunofluorescence analysis of PK-15 cells infected with PSV by laser confocal microscopy. Immunofluorescence assay with DAPI confirmed that the strain SHCM2019 could be recognized by the anti-PSV polyclonal antibody. (D) Transmission electron micrograph (TEM) of cultured PSV strain SHCM2019. Virus pelleted by ultracentrifugation was stained with uranyl acetate and sprayed onto coated formvar grid.
**Figure 2**

Phylogenetic analysis of the polyprotein nucleotide sequence for SHCM2019. A phylogenetic tree was constructed by the neighbour-joining method using the MEGA7.0 software. The numbers at the branches are bootstrap values (%) following 1000 replicates. A dot has been added to the front of the PSV strain isolated in this study. All Chinese isolates in this study were marked in red boxes.
Figure 3

Identification of the recombination events based on the polyprotein gene of the potential recombinants. Similarity analysis of the polyprotein gene of potential recombinants using the SimPlot program with a 200 bp window size, 20 bp step size and 100 bootstrap replicates. The cut-off value in the bootstrapping test (>70%) is indicated by a dotted line, and the arrows indicate putative recombination breakpoints. (A) SHCM2019 represents the query sequence, and HeB04 (red line) and ISU-SHIC (blue line) represent the potential parental strains. (B) Genome origination and recombination-breakpoint identification in SHCM2019. The red area indicates the region of origin from ISU-SHIC, the blue area indicates the region of origin from HeB04, and the white area indicates an unknown origin.
Figure 4

Clinical symptoms and autopsy results of animals. (1) Clinical features around the anus in animals. The piglets in healthy control group were clean and had no attachments around the anus (A). The piglets in PSV-infected group developed watery diarrhoea with residues around the anus (D). (2) Autopsy results of animal intestines. The intestines of piglets in healthy control group were thick and elastic (B). In PSV-infected group, the intestines thinned and its contents were pulpy (E). (3) Autopsy results of animal lungs. The lung performance of piglets in healthy control group was normal (C). Local bleeding points appeared in the lungs of piglets in PSV-infected group (F).
Figure 5

Histopathological examination. (1) Examination of intestine. The intestinal villi of piglets in healthy controls (A). The intestinal villi of pigs infected with PSV were atrophic with obvious oedema of lamina propria and submucosa (C). (2) Examination of lungs. Lung pathology of piglets in healthy controls (B). The alveolar septa of pigs infected with PSV were thickened and had pale serous fluid exudation and many red blood cells in the alveolar septa (D).
Figure 6

Pathogenicity of the SHCM2019 strain in neonatal piglets. (A) Viral RNA shedding in faecal swabs after virus and DMEM inoculation. High levels of viral RNAs were shed in faecal samples of PSV-infected pigs (L1 and L2) at 3–6 dpi, and faecal swabs from DMEM-inoculated piglets (C1 and C2) were PSV viral RNA-negative. (B) IgG antibody levels were measured in infected and control pigs, and antibody levels increased significantly after 6 dpi. (C-D) Viral RNAs were detected in some alimentary tract organs. PSV can be detected in duodenum, jejunum, ileum, caecum, colon, rectum, lung, brain and intestinal lymph, but not in the heart, liver and kidney.

Supplementary Files

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