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Aminopeptidase-N-independent entry of porcine epidemic diarrhea virus into Vero or porcine small intestine epithelial cells

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ABSTRACT

A monkey cell line Vero (ATCC CCL-81) is commonly used for porcine epidemic diarrhea virus (PEDV) propagation in vitro. However, it is still controversial whether the porcine aminopeptidase N (pAPN) counterpart on Vero cells (Vero-APN) confers PEDV entry. We found that endogenous expression of Vero-APN was undetectable in the mRNA and the protein levels in Vero cells. We cloned the partial Vero-APN gene (3340-bp) containing exons 1 to 9 from cellular DNA and subsequently generated two APN-knockout Vero cell lines by CRISPR/Cas9 approach. PEDV infection of two APN-knockout Vero cells had the same efficiency as the Vero cells with or without neuraminidase treatment. A Vero cells stably expressing pAPN did not increase PEDV production. SiRNA-knockdown of pAPN in porcine jejunum epithelial cells had no effects on PEDV infection. The results suggest that there exists an additional cellular receptor on Vero or porcine jejunal cells independent of APN for PEDV entry.

1. Introduction

Porcine epidemic diarrhea virus (PEDV) is a positive-sense single stranded RNA virus belonging to the alphacoronavirus genus in the subfamily Coronavirinae of the family Coronaviridae (Pensaert and de Bouch, 1978; Song and Park, 2012). PEDV causes acute enteric disease in swine, characterized by acute vomiting and watery diarrhea, which has high mortality rates in newborn piglets (Huang et al., 2013; Pan et al., 2012). PEDV was discovered in the United Kingdom in early 1970s and was subsequently identified in many European and Asian countries (Lee et al., 2010; Li et al., 2012; Pensaert and Callebaut, 1974). PEDV field strains isolated before 2010 and the derived vaccine strains belong to genogroup 1 (G1) (Kocherhans et al., 2001). Since late 2010, variant PEDV strains (genogroup 2 [G2]) have emerged in China and Southeast Asia that were fatal to young pigs (Huang et al., 2013; Li et al., 2012). In May 2013, PEDV G2 strains suddenly emerged in the United States, wiping out more than 10% of America's pig population in one year (Huang et al., 2013; Tian et al., 2014). PEDV, together with the other newly emerged swine enteric coronaviruses such as porcine deltacoronavirus (PDCoV) and swine enteric alphacoronavirus (SeAcCoV), are considered serious threats to the pork industry in Asia currently (Jung et al., 2016; Pan et al., 2017).

Aminopeptidase N (APN or CD13) is a type II zinc metalloprotease that mediates various cellular processes, including antigen presentation, cell differentiation, cell motility and coronavirus entry (Luan and Xu, 2007). Porcine APN (pAPN) was first identified as a major receptor for a porcine enteropathogenic alphacoronavirus, transmissible gastroenteritis virus (TGEV) (Delmas et al., 1992). A related human alphacoronavirus, HCoV-229E, utilizes the human APN (hAPN) to enter host cells (Yeager et al., 1992). The interactions between TGEV/HCoV-229E and pAPN/hAPN are highly natural host specific, in which TGEV can use pAPN but not hAPN as its cellular receptor, and HCoV-229E can use hAPN but not pAPN (Kolb et al., 1996). It was reported that transfection and expression of pAPN was sufficient to confer PEDV infection to a non-permissive canine kidney MDCK cell line, indicating pAPN is likely a functional receptor for PEDV as well (Li et al., 2007). Moreover, transgenic mice expressing pAPN confer susceptibility to PEDV (Park et al., 2015). It was also reported that, unlike TGEV or HCoV-229E, PEDV spike protein-mediated pseudovirus was able to enter non-permissive canine kidney MDCK cells exogenously expressing either hAPN or pAPN, suggesting that the PEDV-APN interaction is not species specific (Liu et al., 2015). Correspondingly, PEDV can infect various cell lines from human, monkey, pig and bat species (Liu et al., 2015), but whether species-specific APN functions as the PEDV entry receptor on the respective cell line has not been investigated. However, most recently, two groups independently demonstrated that neither hAPN nor pAPN is the functional receptor for PEDV by using more comprehensive analyses, including knockout of endogenous expression of hAPN or pAPN in human or porcine cell lines by the Clustered Regularly Interspaced Short Palindromic Repeats (CRISPR)/CRISPR-
associated protein-9 nuclease (Cas9) system (Li et al., 2017; Shirato et al., 2016).

An African green monkey (Chlorocebus sabaeus) kidney cell line, Vero (ATCC CCL-81), is commonly used for PEDV adaption and culture in the presence of trypsin in vitro for research or vaccine production (Hofmann and Wyler, 1988; Pan et al., 2012). Green monkey specific APN expressed on Vero cells (Vero-APN/vAPN) may serve as the entry receptor for PEDV (Li et al., 2007). However, a few studies remarked Vero cells without APN expression, but no experimental evidences were provided in these publications (Li et al., 2016; Nam and Lee, 2010). Therefore, the aim of this study was to clarify the argument, investigating whether Vero-APN is indeed responsible for PEDV entry into Vero cells. Furthermore, we sought to test whether knockdown of pAPN expression on porcine small intestinal epithelial cell line, IPEC-J2, is correlated with PEDV infection in comparison with TGEV.

2. Results and discussion

We first aimed to confirm if the putative green monkey APN (vAPN) gene exists in Vero cellular genome DNA. The 5′-part of the vAPN gene (3,340 bp), which was not available on the public genome database, was cloned by genomic PCR from extracted Vero genomic DNA with a pair of PCR primers shared significant sequence homology with human, chimpanzee, rhesus monkey and pig APNs. As illustrated in Fig. 1, the partial cloned vAPN gene contains nine exons or coding DNA sequence (CDS). The start codon ATG is located at CDS1 having 617 base pair (bp), whereas the sizes of the other eight CDS (CDS2–9) vary from 66 to 155 bp (Fig. 1). The partial cloned vAPN gene and its encoding cDNA (1,572 bp in size) share 97.9% and 98.9% sequence homology with the available rhesus monkey APN gene and cDNA, respectively. Attempts to amplification of the entire or the 5′-partial vAPN mRNA by RT-PCR were failed. Therefore, we performed a quantitative real-time PCR (qRT-PCR) analysis to detect if vAPN mRNA is expressed endogenously in Vero cells. As a control we examined hAPN mRNA expression in Huh-7 cells. When we confirmed endogenous hAPN mRNA expression, which was in line with the previous publications (Li et al., 2017; Liu et al., 2015), relative quantification of vAPN mRNA level to the green monkey β-actin control in Vero cells did not give a convincingly positive value in a cell-culture period of 3 days (data not shown). The result indicated that vAPN mRNA was expressed under the detection limit or was deficient in expression.

We further examined endogenous and exogenous expression of APN in the protein level using a broadly reactive anti-APN antibody by immunofluorescence assay (IFA) and western blot (WB) analysis. For comparison, two recombinant APN expression plasmids were constructed. Plasmid pCI-pAPN harbored the full-length porcine APN cDNA (2,892 bp) whereas plasmid pRK-vAPN contained the nine CDS of vAPN without introns followed by a stop codon (1,575 bp) that were fused sequentially by overlapping PCR. Vero or Huh-7 cells were transfected with either of the two plasmids. At 48 hour post-transfection, transfected or untransfected cells were subjected to IFA or WB. Specific anti-APN fluorescence signal was detected in transfected Vero or Huh-7 cells as well as untransfected Huh-7 cells (though the intensity was weak), but not observed in untransfected Vero cells (Fig. 2A). The WB analysis was consistent with the IFA result, showing individual bands of distinct sizes representing the full-length APN (APN-FL) or the partial APN (vAPN-CDS1–9) in pCI-pAPN- or pRK-vAPN-transfected Vero cells, whereas no bands were detected in untransfected Vero cells (Fig. 2B). In contrast, Huh-7 cells displayed a band corresponding to APN-FL regardless whether transfection with pCI-pAPN- or pRK-vAPN or not (Fig. 2B). These results indicated that Huh-7 cells express hAPN endogenously while Vero cells probably do not express vAPN inherently.

To further examine the effects of anti-APN antibody on PEDV entry into these two cell lines, we carried out quantitative PEDV infection analysis with a recombinant PEDV-GFP (see “Materials and methods”). Either Vero or Huh-7 cells were infected efficiently by PEDV-GFP, showing clear GFP fluorescence co-localized with cytopathic effects (CPE), which could be assessed by counting the numbers of GFP-positive cell clusters (Fig. 3A). Moreover, PEDV-GFP infection of two cell lines could be neutralized by anti-PEDV-S IgM or IgG in a dose-dependent manner (Fig. 3B), indicating that it is an appropriate in vitro model for quantification of PEDV infection. When anti-APN antibody at 2 μg/ml or 20 mg/ml was incubated with the cells prior to PEDV-GFP infection, it did not block PEDV-GFP entry or infection in Vero cells at 12 or 24 hour post-infection (hpi; Fig. 3C). For Huh-7 cells, it was observed that pretreatment with the antibody significantly decreased GFP-positive cell cluster numbers at 12 hpi; however, there was no differences in comparison with untreated cells at 24 hpi (Fig. 3C). Shirato and his colleagues reported that pAPN promotes PEDV infection through its protease activity rather than its proposed receptor function (Shirato et al., 2016). Since the protease active sites are highly conserved between hAPN and pAPN (Delmas et al., 1994), it is hypothesized that the anti-APN antibody blocked PEDV infection by targeting the enzymatically catalytic sites of hAPN on Huh-7 cells in the early stage (0–12 hpi). The inhibited effect was likely diminished when PEDV continued to propagate and spread thereafter. These results demonstrated that anti-APN antibody had no effects on PEDV entry into Vero cells, probably due to the absence of vAPN expression.

In order to completely rule out the potential vAPN production in Vero cells that may be under the detection limit, we set out to generate vAPN-knockout cells using CRISPR/Cas9 system. A single-plasmid CRISPR/Cas9 approach was developed, in which two guide RNA (gRNA) molecules and the Cas9 enzyme followed by a puromycin resistance gene are simultaneously expressed (Fig. 4A). We designed a knockout strategy using an engineered pX480-vAPNKO plasmid by targeting the vAPN CDS1 with two gRNA/Cas9 complexes to delete a 160-bp fragment between nucleotides 290–449 downstream of the translation initiation site (Fig. 1). At 3 days post-transfection with the pX480-vAPNKO plasmid plus puromycin selection, the genomic DNA was isolated and tested for vAPN gene integrity by PCR. It was shown that two different cell pools (using the same approach in two batches) possessed the intended vAPN deletion fragment, displaying weak bands smaller than the intact CDS1 (Fig. 4B). From these cell pools, we generated single cell clones, designated as vAPNKO1 and vAPNKO2, by limiting dilution with puromycin selection again. However, the intended vAPN deletion fragment was not visible in the clone vAPNKO1 (Fig. 4C). Sequence analysis of the PCR products of the clone vAPNKO1 revealed that there was only a 5-bp deletion (nt 290–294) downstream of the first Cas9 cleavage site (Fig. 4D). The single cell clone vAPNKO2 displayed the intended vAPN deletion fragment that was subsequently confirmed to have the expected 160-bp deletion by sequencing (Fig. 4C and D). Nevertheless, either of the deletions in vAPNKO1 or vAPNKO2 would result in a frameshift of vAPN allele.

We next confirmed that these knockout cells, Vero-vAPNKO1 and Vero-vAPNKO2, were deficient in vAPN expression by IFA (Fig. 5A). Upon PEDV-GFP infection, both Vero-vAPNKO1 and Vero-vAPNKO2 cells displayed the intended vAPN deletion fragment that was subsequently determined in this study (GenBank accession number KX342858). Arrowheads indicate two Cas9 cleavage sites (nucleotides 290–449) downstream of the translation initiation site (ATG) within the exon 1 (CDS1). Deletion of a 160-bp fragment between C.-M. Ji et al. Virulence 517 (2018) 16–23
exhibited consequent CPE and GFP expression similar to normal Vero cells, indicating that vAPN knockout had no effects on PEDV entry and infection (Fig. 5B). Sialic acids such as Neu5Ac have been shown to be involved in PEDV entry as the co-receptor (Liu et al., 2015); therefore we examined whether there may be any interplay between sialic acids and APN for PEDV entry. Vero, Vero-vAPNKO1 and Vero-vAPNKO2 cells were infected with PEDV-GFP, respectively, in the presence of distinct concentrations of neuraminidase, which would cleave sialic acids on the cell surface. PEDV-GFP infection was inhibited slightly by neuraminidase treatment in a dose-dependent manner (Fig. 5C). However, there were no significant differences of overall effect of neuraminidase treatment on PEDV-GFP infection between normal Vero cells and knockout cells (Fig. 5C). These results indicated that vAPN knockout had no effects on potential association with sialic acids mediating PEDV entry.

It is reported that overexpression of pAPN in porcine kidney or testis cells slightly increased the production of PEDV, suggesting the role of pAPN in promoting PEDV replication (Nam and Lee, 2010; Shirato et al., 2016). We developed a stable Vero cell line expressing pAPN (Vero-pAPN; (Wang et al., 2018), and then determined whether Vero-pAPN cells slightly increased the production of PEDV, suggesting the role of entry.

Wild-type PEDV-GFP was used to infect two cell lines (36–48 hpi; Fig. 6). The result indicated that although pAPN slightly facilitated PEDV replication in the early stage, it had limited effects on PEDV infection in Vero cells, probably due to the presence of an unidentified bona fide receptor.

PEDV infects pigs by targeting porcine small intestinal epithelial cells. Recently, a non-transformed, non-tumorigenic cell line named IPEC-J2, from jejunum epithelium isolated from a neonatal piglet, was characterized and used for studying porcine enteric virus-host interactions (Liu et al., 2010). Efficient PEDV-GFP or TGEV infection of IPEC-J2 cells was confirmed by optimizing the multiplicity of infection even treated with a small interfering RNA (siRNA) negative control (Fig. 7A and B), allowing us to use this cell line to study PEDV-pAPN interplay. However, our many attempts to knockout the pAPN in IPEC-J2 cells by CRISPR/Cas9 technique to knockout either hAPN in Huh-7 cells or vAPN in ST cells, still resulting in efficient infection of PEDV in knockout cells, which convincingly demonstrated that PEDV infection does not utilize them as functional receptors (Li et al., 2017; Shirato et al., 2016). In particular, Li et al. employed CRISPR/Cas9 technique to knock out hAPN in Huh-7 cells or pAPN in ST cells, still resulting in efficient infection of PEDV in knockout cells, which convincingly demonstrated that PEDV infection does not utilize them as functional receptors (Li et al., 2017). Our study is in line with these results, providing additional evidences that the Vero cell line of green monkey origin does not express vAPN for PEDV entry and propagation, and that PEDV infection of the small intestinal epithelial cells is likely not associated with pAPN. The present study
should have broad biological significance, since the Vero cell is commonly used for PEDV propagation and vaccine production in vitro, and the porcine small intestine epithelial cells is the PEDV target in vivo. Furthermore, in our experiment using the PEDV non-susceptible BHK-21 or NIH-3T3 cells, we found that overexpression of pAPN did not confer PEDV-GFP entry or infection as well (data not shown). Therefore, these data collectively suggest that there exist additional cellular receptors independent of human, green monkey, and porcine APN for PEDV entry. Moreover, although PEDV is able to infect various cell lines of distinct species origin (Liu et al., 2015), even in duck species (Khatri, 2015), the other non-porcine and non-primate species-specific APNs (such as bat or duck APN) likely not functions as the PEDV entry receptor. Searching for the unknown PEDV functional receptor is still ongoing and should answer this enigma in the future.

3. Materials and methods

3.1. Cell lines and virus stocks

A monkey kidney cell line Vero (ATCC CCL-81), human hepatoma cell line Huh-7 and a swine testis cell line ST (ATCC CRL-1746) were grown in Dulbecco’s modified Eagle medium (DMEM) supplemented with 10% fetal bovine serum (FBS) and 1% antibiotics at a 37°C incubator, respectively. A Vero cell line stably expressing the pAPN coding region (Vero-pAPN) was cultured in DMEM supplemented with 10μg/ml puromycin and antibiotics (Wang et al., 2018). The IPEC-J2 cell line, from jejunum epithelium isolated from a neonatal piglet, was a generous gift from Dr. Lijuan Yuan at Virginia Tech, Blacksburg, VA (Liu et al., 2010). The IPEC-J2 cells were grown in DMEM supplemented with 5% FBS and 1% antibiotics.

A G2 PEDV Chinese strain, ZJU/G2/2013 (GenBank accession no. KU558701), was used in this study (Qin et al., 2017). The virus stocks of recombinant ZJU/G2/2013 expressing enhanced green fluorescent protein (GFP), designated PEDV-GFP, was generated by transfection of Vero cells with a DNA-launched PEDV infectious cDNA clone (Zhao et al., 2018), followed by two serial passages on Vero cells supplemented with 5μg/ml trypsin (Sigma, Cat#T7186-50TAB, St Louis, MO, USA) and without FBS. The virus stocks were PEDV-GFP-containing supernatants without cell debris that were removed by centrifugation. The virus titers of PEDV-GFP were determined by endpoint dilutions as 50% tissue culture infective dose (TCID50) on Vero cells. Virus stocks were stored at −80 °C until use. The PEDV-GFP infection efficiency in Vero, IPEC-J2, or Huh-7 cells was assessed by counting the numbers of GFP-positive cell clusters (quantified as fluorescent focus-forming unit [FFU]). TGEV Purdue strain was kindly provided by Dr. Rong Ye at Shanghai Medical College of Fudan University, which was propagated on ST cells (Pan et al., 2017). The TGEV infection efficiency in TGEV-infected-IPEC-J2 cells was also evaluated by counting the FFU numbers following immunofluorescence assay with a TGEV-specific antibody.

3.2. Genomic PCR and sequence analysis

A pairs of primers, νAPN-CDS1-F (5’: ATGGCCAAGGGCTTCTACAT TTCAGG-3’) and νAPN-CDS9-R (5’: CTCTCGAGGTGTCCCACAGG TTC-3’), were designed for amplification of the S’-end region of the putative green monkey APN gene in Vero cells by one-step genomic
PCR, based upon four published APN gene sequences from different species: *Homo sapiens* (human; Gene ID: 290), *Pan troglodytes* (chimpanzee; Gene ID: 467758), *Macaca mulatta* (rhesus monkey; Gene ID: 701117) and *Sus scrofa* (pig; Gene ID: 397520). The forward primer vAPN-CDS1-F contains the start codon ATG while the reverse primer vAPN-CDS9-R is complementary to the sequence at the 3'-end of the exon-9 of the rhesus monkey APN gene. Genomic PCR was performed with a KOD DNA polymerase (Toyobo, Japan) using 100 ng of extracted genomic DNA from Vero cells in a total volume of 50 μl. The PCR condition was 30 cycles of 98°C for 30 sec, 58°C for 30 sec, 68°C for 4 min with an initial denaturing of the template DNA at 94°C for 3 min and a final extension at 68°C for 7 min. The resulting fragment was cloned into a Zero pCR-Blunt vector (Thermo Fisher Scientific) by blunt-end cloning strategy followed by Sanger sequencing. Analysis and alignment of DNA and amino acid sequences were performed using Lasergene Package (DNASTAR Inc., Madison, WI).

3.3. Construction of the recombinant vectors expressing pAPN or Vero-APN and in vitro expression

The complete coding region of pAPN was amplified by one-step RT-PCR with primers PCI-APN-F (aggctagctACCATGGCCAAGGGATTCTA CATTTCCA) and PCI-APN-R (TTGTGACCTAGCCTGTGCTATGAAC CAATTCAACA) using total RNAs extracted from porcine small intestine, and subsequently cloned into a pCI-neo vector (Promega) downstream of the CMV immediate-early enhancer/promoter using NheI and SalI restriction sites. The construct was sequenced to verify the identity and designated as pCI-pAPN. The nine exons, encoding the first 1572-bp of Vero-APN mRNA from the Vero-APN gene, were fused sequentially by overlapping PCR, and inserted into a pRK5 vector, to construct an expression vector, pRK-vAPN. Plasmids pCI-pAPN or pRK-vAPN were transiently transfected into Vero cells using Lipofectamine 3000 (Thermo Fisher Scientific) according to the manufacturer’s protocol. Cells were cultured for 48 to 72 hours, and then applied to immunofluorescence assay or western blot to detect the expression of APN.

3.4. siRNA and siRNA transfections

The siRNA directed against pAPN (sipAPN: 5'-AACAAGCCCAGCU GGUAAA-3') or the negative control (siNC: 5'-UUCUCCGAACGUGUC AC GU-3') was purchased from RiboBio (Guangzhou, China). The siRNA transfections in IPEC-J2 cells were performed using the Lipofectamine RNAiMAX transfection reagent (Thermo Fisher Scientific, USA) according to manufacturer's instructions. At 24 hour post-transfection, IPEC-J2 cells were infected with PEDV-GFP at an MOI (multiplicity of infection) of 5 or with TGEV (MOI = 1).

3.5. Immunofluorescence assay, western blot and quantitative real-time PCR

Transfected cells or APN-knockout Vero cells were washed 2 times with PBS, fixed with 4% paraformaldehyde in PBS for 20 min and then
permeabilized with 0.5% Triton X-100 for 10 min. One hundred microliters of the polyclonal anti-APN antibody (abcam #93897), at a 1:100 dilution in PBS, was added over the cells and incubated for 1 h at 37°C. Cells were washed twice with PBS and 100 μl FITC-labeled goat anti-rabbit IgG (Thermo Fisher Scientific) at a 1:100 dilution was then added. After 30 min incubation at 37°C, the cells were washed trice with PBS and 100 μl 1% Triton X-100 for 10 min. One hundred microliters of the polyclonal anti-APN antibody at a 1:200 dilution followed by incubation with horse-radish peroxidase (HRP)-conjugated anti-rabbit IgG (Thermo Fisher Scientific), Detection of the N protein expression of PEDV or TGEV in infected IPEC-J2 cells were conducted by using the anti-PEDV-N monoclonal antibody (purchased from Medgene Labs, Brookings, SD, USA) or the anti-TGEV-N polyclonal antibody.

For qRT-PCR analysis, total RNA was extracted from cell lysing using an AxyPrep Multisource Total RNA Miniprep Kit (Axygen). Detection of mRNA expression levels of hAPN in Huh-7 cells or vAPN in Vero cells by qRT-PCR was performed using TransScript Green One-Step qRT-PCR SuperMix (Transbio, Beijing, China) with primers hAPN-qPCR-F (5'-GGACAGCCAGTATGAGAT-3') and hAPN-qPCR-R (5'-GGATAACGG TGATGTTGAA-3'), or primers vAPN-qPCR-F (5'-TGAGATGGCAGTAGTA GTTCC-3') and vAPN-qPCR-R (5'-GTGGATAAGGCTGTTGGT-3'). Relative quantification was expressed as $2^{-ΔΔCt}$, where $ΔCt$ is the difference between the main Ct value of the sample in triplicate and that of an endogenous human β-actin (primers 5'-ATGGAGTGTACACCCGTGCTCC TCC-3' and 5'-GGAATCAAAGAATTGGTGGTTC-3') or green monkey β-actin control (primers 5'-CTGGGGAGCGAGGTCTGAG-3' and 5'-GGGCCCC GACCTGTCATAC-3').

### 3.6. Treatment with anti-PEDV spike (S) protein antibodies, anti-APN antibody or neuraminidase

Cells were incubated with the anti-APN antibody at 2 μg/ml or 20 mg/ml or neuraminidase (Sigma) at concentrations of 5, 10, 20, 30 or 50 μU in DMEM for 1 h at 37°C to prevent PEDV-GFP infection. For neutralizing test, serial dilutions (1:50, 1:100, 1:500 or 1:1000) of anti-PEDV-S IgM (Medgene Labs, Brookings, SD, USA) or anti-PEDV-S IgG (JBT, Korea) were each mixed with 100 FFU of PEDV-GFP, incubated at 37°C for 1 h to form virus-antibody complexes, and added to Vero cells. Cells were fixed in 4% parafomaldehyde at appropriate time points, and GFP-positive cells were counted.

### 3.7. Generation of APN knockout (KO) cell lines by using CRISPR/Cas9 system

We first constructed a Cas9 backbone cloning vector, named pX480, harboring two guide RNA (gRNA) expression cassettes as well as a 2A-puromycin resistance gene inserted following the human Cas9 gene based upon the pX330 vector (Addgene) (Ran et al., 2013). The used primers and construction details will be available upon request. The pX480 vector was digested with BbsI and BsaI sequentially and ligated polystyrene cell cultures (PLC) membrane that was subsequently blocked with Tris-buffered saline (TBS) containing 3% bovine serum albumin (BSA) overnight at 4°C. Proteins were detected using the anti-APN antibody at a 1:200 dilution followed by incubation with horse-radish peroxidase (HRP)-conjugated anti-rabbit IgG (Thermo Fisher Scientific).
with annealed protospacer oligoDNAs (nucleotides 284–306, 5'-CCGA TGACAGGGGCCTATAC-3' and nucleotides 444–466, 5'-CGACATCGAC AGAACCGAGC-3') specific for the vAPN CDS1 sequences to obtain a single-plasmid pX480-vAPNKO. Vero-APN knockout cells were generated by CRISPR/Cas9 gene editing as described (Ran et al., 2013). Single cell clones of vAPN knockout cells were obtained by limiting dilution and genotyped by PCR and DNA sequencing.

3.8. Nucleotide sequence accession numbers

The sequences of the pAPN cDNA and the partial Vero-APN gene determined in this study have been deposited in GenBank under accession nos. KX342854 and KX342855, respectively.

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Fig. 7. Effects of pAPN knockdown on PEDV-GFP or TGEV infection in IPEC-J2 cells. (A) IPEC-J2 cells were transfected with sipAPN or siNC for 24 hours followed by infection with either PEDV-GFP (MOI = 5) or TGEV (MOI = 1). Observation of GFP expression in PEDV-GFP-infected cells or detection of TGEV-N protein expression in TGEV-infected cells by IFA was performed at 24 hpi. (B) The numbers of PEDV- and TGEV-positive cell clusters were measured from (A), respectively. The result represented mean values from three independent experiments, and error bars indicate standard deviation (***, p < 0.001). (C) Detection of expression of pAPN, TGEV-N or PEDV-N in siRNA-treated cells followed by virus infection (24 hpi) by western blotting.
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