Mutation Profiles of eGFP-Tagged Small Ruminant Morbillivirus During 45 Serial Passages in Ribavirin-Treated Cells

Fuxiao Liu1†, Yanli Zou2†, Lin Li2, Chunju Liu2 and Xiaodong Wu2*

1 College of Veterinary Medicine, Qingdao Agricultural University, Qingdao, China, 2 OIE Reference Laboratory for Peste des Petits Ruminants, China Animal Health and Epidemiology Center, Qingdao, China

Small ruminant morbillivirus (SRMV), formerly known as peste-des-petits-ruminants virus, classified into the genus Morbillivirus in the family Paramyxoviridae. Its L protein functions as the RNA-dependent RNA polymerases (RdRp) during viral replication. Due to the absence of efficient proofreading activity in their RdRps, various RNA viruses reveal high mutation frequencies, making them evolve rapidly during serial passages in cells, especially treated with a certain mutagen, like ribavirin. We have previously rescued a recombinant enhanced green fluorescence protein-tagged SRMV (rSRMV-eGFP) using reverse genetics. In this study, the rSRMV-eGFP was subjected to serial passages in ribavirin-treated cells. Due to the ribavirin-exerted selective pressure, it was speculated that viral progenies would form quasispecies after dozens of passages. Viral progenies at passage-10, -20, -30, -40, and -50 were separately subjected to next-generation sequencing (NGS), consequently revealing a total of 34 single-nucleotide variations, including five synonymous, 21 missense, and one non-sense mutations. The L sequence was found to harbor eight missense mutations during serial passaging. It was speculated that at least one high-fidelity variant was present in viral quasispecies at passage-50. If purified from the population of viral progenies, this putative variant would contribute to clarifying a molecular mechanism in viral high-fidelity replication in vitro.

Keywords: rSRMV-eGFP, next-generation sequencing, single-nucleotide variation, mutation, L protein, high-fidelity

INTRODUCTION

Peste des petits ruminants is a highly contagious disease, mainly affecting goats and sheep, and even large ruminants (1–3). Its etiological agent is small ruminant morbillivirus (SRMV), formerly known as peste-des-petits-ruminants virus. SRMV belongs to the genus Morbillivirus in the family Paramyxoviridae. Its genome is a single strand of RNA with negative polarity (4), coding for six structural proteins, namely nucleocapsid (N), phospho (P), matrix (M), fusion (F), hemagglutinin (H), and large (L) proteins in the order of 3′-N-P (V/C)-M-F-H-L-5′. The V and C proteins are two non-structural proteins, encoded through alternative expression strategies from the P gene transcription unit (5). Viral RNA genome is encapsulated by the N protein forming a helical nucleocapsid, in combination with the L and P proteins to form a ribonucleoprotein complex (6). The L protein is an RNA-dependent RNA polymerase (RdRp),
the largest of the morbilliviral proteins, and is least abundant in virus-infected cells. It is assumed to carry all activities necessary for genomic RNA transcription and replication (5).

The overwhelming majority of RNA viruses use their own low-fidelity RdRp to initiate RNA synthesis (7–9), consequently generating viral quasispecies consisting of a swarm of viruses. Compared with most DNA viruses, morbilliviruses are characterized by high mutation frequencies in their genomes and antigenomes during viral serial passages in vitro, mainly attributed to the lack of effective proofreading activities in their RdRps (10). For example, the measles morbillivirus was estimated to have a mutation rate of 9 × 10^{-5}/nt/replication and a genomic mutation rate of 1.43/replication by analysis on the frequency of monoclonal antibody-resistant mutants (11). More recently, Zhang et al. (12) determined a spontaneous mutation rate of 1.8 × 10^{-5}/nt/replication in measles morbillivirus under non-selective conditions.

Using Sanger sequencing, we have reported major mutation events in structural genes of a China SRMV isolate (China/Tibet/Geg/07-30, GenBank: FJ905304.1) during 90 serial passages in vitro (13). However, the conventional Sanger sequencing is not capable of generating a large dataset to identify all single-nucleotide variations (SNVs) in viral progenies. As an alternative technique, the next-generation sequencing (NGS) has been broadly applied to analyze and to quantify the exceptionally-high diversity within viral quasispecies. By means of it, single-nucleotide polymorphisms (SNPs) in a variety of viruses can be systematically analyzed to reveal their evolutionary profiles (14–18).

Enhanced green fluorescence protein (eGFP) is an ideal marker, which facilitates screening of high-fidelity mutants to unveil a given mechanism concerning high-fidelity replication of RNA virus (19). We have recovered an eGFP-tagged recombinant SRMV (rSRMV-eGFP) from the cDNA clone of Nigeria 75/1 vaccine strain (20). In the present study, the rSRMV-eGFP was subjected to 45 serial passages in Vero-Dog-SLAM (VDS) cells. In order to quicken the process of SNVs in the rSRMV-eGFP genome, VDS cells were always treated with ribavirin, a widely used mutagen that could force RNA viruses into mutagenesis, and even into error catastrophe (21, 22). The NGS technique was used to uncover the mutation profiles of viral progenies at different passages.

**MATERIALS AND METHODS**

**Cell and Virus**

The VDS cell line was cultured at 37°C with 5% CO2 in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (FBS), and containing penicillin (100 U/mL), streptomycin (100 µg/mL), amphotericin B (0.25 µg/mL), and G418 (500 µg/mL). The rSRMV-eGFP was rescued previously in our laboratory (20).

**Forty-Five Serial Passages of rSRMV-eGFP**

The passage-5 (P5) rSRMV-eGFP was serially passaged (3 d/passage) in ribavirin (Solarbio, Beijing, China)-treated VDS cells at undetermined MOI values for 45 passages (P6 to P50) in the biosafety level-3 laboratory of China Animal Health and Epidemiology Center. The ribavirin concentration was gradually increased in DMEM with passaging: 80 µM during P6–16, 160 µM during P17–24, 240 µM during P25–32, 320 µM during P33–42, and 400 µM during P43–50. Virus-infected cells were observed under a fluorescence microscope before each round of passaging.

**NGS Analyses of P10, P20, P30, P40, and P50 Progenies**

Culture supernatants with rSRMV-eGFP-infected cells were independently harvested from the P10, P20, P30, P40, and P50 progenies for extracting viral RNA using the Viral RNA/DNA Extraction Kit (Takara, Dalian, China), according to the manufacturer’s instruction. A total of five RNA samples were separately reverse transcribed using the 1st Strand cDNA Synthesis Kit (Takara, Dalian, China) and random hexamers, according to the manufacturer’s instruction. The Illumina sequencing and library construction were performed as described previously (18). In brief, the NEBNext® Ultra™ II RNA Library Prep Kit (NEB, Ipswich, MA, USA) was used for library construction. After adapter ligation, 10 cycles of PCR amplification were performed for sequencing target enrichment. The libraries were pooled at equal molar ratio, denatured, and diluted to optimal concentration prior to sequencing. The Illumina NovaSeq 6000 (Illumina, San Diego, CA, USA) was used for sequencing to generate pair-end 150 bp reads.

**Processing and Analysis of NGS Data**

Processing and analysis of NGS data were also carried out as described previously (18). In brief, raw reads were filtered by fastp (https://github.com/OpenGene/fastp) to remove sequencing adapters and low-quality reads, including those reads scored < Q20. Ribosomal RNAs and host reads subtraction by read-mapping were performed with BBMap program (https://github.com/BioInfoTools/BBMap). A de-novo genome assembly was performed using SPAdes v3.14.1 (https://github.com/ablab/spades). These extracted assembled scaffolds limited the minimum contig length to 100 bases, with the best BLAST hits to the NCBI nucleotide database. High-quality filtered reads were mapped against the rSRMV-eGFP reference genome by Burrows-Wheeler Aligner v0.7.17 (http://bio-bwa.sourceforge.net/), which also generated a BAM file to calculate the mapping depth and coverage. SNPs were identified using an integrated software package, Snippy v4.4.5 (https://github.com/teemann/snippy), which included both substitutions and insertions/deletions. The available SNP results were selected if mapping quality was ≥ 60 and depth was ≥ 10.

**RESULTS**

**Fluorescence-Attenuated or -Disappeared Syncytia Become Visible With Passaging**

The P5 rSRMV-eGFP was serially passaged in ribavirin-treated VDS cells. Fluorescent syncytia were always observable during passaging (Figure 1). However, a few fluorescence-attenuated syncytia occasionally appeared on cell monolayers approximately
after the 24th passaging (Figure 1, enclosed by yellow line at P31), implying that the eGFP open reading frame (ORF) was undergoing missense mutations, or nucleotide deletions from the viral genome. A few fluorescence-disappeared syncytia (Figure 1, enclosed by white line at P43) became visible at P43 but were uneasily found on the virus-inoculated cell monolayer at 72 h post-infection (hpi). The proportion of fluorescence-disappeared syncytia increased with passaging and even was predominant in a randomly selected field-of-view of fluorescence microscope at P50 (Figure 1, enclosed by white line).

**NGS Shows Analyzable Sequencing Depths**

The rSRMV-eGFP had a 16788-nt-long genome. Figure 2A schematically showed all ORFs and untranslated regions (UTRs) in proportion to their actual distributions in the viral antigenome. To uncover viral mutation profiles with passaging, a total of five RNA samples were independently extracted from the P10, P20, P30, P40, and P50 culture supernatants, and then subjected to the NGS analysis. The NGS data were deeply analyzed to yield acceptable sequencing depths for the P10, P20, P30, and P40, whereas a short GC-rich region between the M and F ORFs showed the lowest depths for all five samples (Figure 2B).

**Thirty-Four SNVs Are Identified in Five Progenies**

The NGS analyses showed that neither sequence-deleting nor inserting phenotype was detectable in the five progenies, within which a total of 34 SNVs arose with passaging, containing 26 transitions and eight transversions. The positions of SNVs were independently indicated with blue arrows in the rSRMV-eGFP antigenome, in which neither ORFs nor UTRs proportionally matched their actual lengths (Figure 3A). Figure 3B revealed complete profiles of 34 SNVs during viral serial passages. Each SNV was identified as a SNP with mixture of two nucleotides. The 5' UTR exhibited one SNV (C28T) only at P50. The other two UTRs, between the M and F ORFs as well as between the F and H ORFs, exhibited five and one SNVs, respectively. Except the C28T and C6094T at P50, the other SNVs showed low frequencies of occurrence in UTRs. Seven ORFs, namely the N, P, eGFP, M, F, H, and L, showed one, three, four, four, three, four, and eight SNVs, respectively. Several SNVs, such as T2469A and T5251C, appeared at a given passage, but reverted back to their original statuses with passaging. In contrast, the site at nt 8184 underwent gradual substitution of a nucleotide for another one with passaging. Its original nucleotide was “A,” which was gradually and finally replaced by “G” at P40.

![FIGURE 1](https://example.com/figure1.png) | Fluorescence-attenuated or -disappeared syncytia formation during 45 serial passages in ribavirin-treated VDS cells. Fluorescence-attenuated and -disappeared syncytia are enclosed by yellow and white lines, respectively. BF, bright field.
were some low-frequency point mutations, like G5948A and G6081A, occurring once only at a single passage.

**Synonymous, Missense, and Non-Sense Mutations Are Identified in Antigenome**

Seven ORFs totally showed 27 of their own SNVs, including five synonymous, 21 missense, and one non-sense mutations. Accordingly, 21 missense mutations caused 21 amino acid (aa) mutations (Figure 3A, black-marked in black brackets), distributed in seven protein sequences. The L protein showed the most aa mutations, up to eight. In the L protein sequence, the SNV “A12813C” occurred at P10 and P20, and its adjacent SNV, “T12814G,” arose at P40. The “A12813C” and “T12814G” led to substitutions of leucine and serine for the same isoleucine at aa 896, respectively. As a non-self-sequence, the eGFP ORF harbored a non-sense mutation at nt 3794, causing the early termination of translation at aa 116 in the eGFP sequence (Figure 3A, red-marked in red bracket). The SNV “G3794T” was initially identified at P40, when it only showed a low frequency of occurrence. Nevertheless, this SNV became predominant in viral quasispecies at P50 (Figure 3B), directly responsible for a number of fluorescence-disappeared syncytia that appeared on the P50 cell monolayer at 72 hpi.

**DISCUSSION**

The SRMV Nigeria 75/1 strain has been widely used for vaccine production. Its reverse genetics system had been constructed previously in our laboratory for further use to rescue recombinant SRMVs expressing foreign proteins (20, 23, 24). The rSRMV-eGFP was demonstrated to have a similar growth curve to that of the wide-type strain (20), implying no significant interference of the eGFP with viral replication. Moreover, the eGFP facilitated observation of fluorescent syncytium formation on a cell monolayer in real time. Such a feature prompted us to use the rSRMV-eGFP for uncovering viral mutation profiles during serial dozens of passages. To promote occurrence of SNVs in the eGFP ORF, VDS cells were treated with the ribavirin, which could act as a viral mutagen forcing RNA viruses into mutagenesis. In order to make the rSRMV-eGFP gradually adapt to the ribavirin-treated cells, the ribavirin concentration was progressively increased with viral passaging. The rSRMV-eGFP was subjected to 45 serial passages in ribavirin-treated cells.
FIGURE 3 | Progeny profiles of nucleotide and amino acid mutations at P10, P20, P30, P40, and P50. All detectable single-nucleotide and -amino acid mutations in rSRMV-eGFP antigenome during passaging (A). The positions of SNVs are independently indicated with blue arrows. SNVs are marked with blue letters and numbers. Missense mutation-caused amino acid substitutions are marked with black letters and numbers in black brackets. The non-sense mutation, G3794T, causes early termination of eGFP translation, marked with “(E116*)” in red. All elements do not match their actual lengths in proportion. Relative depths of next-generation sequencing for 34 SNVs in rSRMV-eGFP antigenome with 45 serial passages (B). The relative depth is referred to as the proportion of sequencing depth of a given nucleotide to the total.
and speculated to form a rich diversity of viral quasispecies at P50, because most RNA viruses are genetically unstable during replication (8).

To explore SNV profiles of the rSRMV-eGFP during serial passages, the P10, P20, P30, P40, and P50 progenies were collected for the NGS analysis. The major reason why the conventional Sanger sequencing was not used in this study was its limited ability or inability to identify low-frequency mutations, seriously reducing its applicability to quantify the complexity of mutant spectra. In contrast, the NGS, because capable of generating a large dataset for identification of SNMs in viral genomes, has been widely used to analyze quantitatively the rich diversity of viral quasispecies (18, 25, 26).

Viral self-proteins are intrinsically encoded in rSRMV-eGFP-infected cells. In order to avoid the impact of an error catastrophe on virus propagation, harmful or especially lethal SNVs would not accumulate in viral self-sequences. Indeed, the NGS analyses revealed no lethal SNV in the viral self-sequences. In contrast, the SNVs were unrestricted in the eGFP ORF during viral propagation, due to the eGFP as a foreign protein, theoretically uninvolved in a series of SRMV-related events, for example, replication, transcription, regulation, and packaging. In other words, SNVs would be random, uncontrolled, and retainable in the eGFP ORF during viral propagation. Indeed, compared with the other ORFs in the rSRMV-eGFP antigenome, the eGFP sequence revealed a high mutation frequency, as evidenced by the NGS analyses.

A total of four SNVs, including one non-sense mutation, appeared in the eGFP ORF during serial passages. The first one (C3641A) was found to arise at P30, causing substitution of methionine for leucine at aa 65. It remained unclear whether the “L65M” resulted in the fluorescence-attenuated syncytium formation on cell monolayer at P31 (Figure 1, P31). The unique non-sense mutation arose in the eGFP ORF at P40, directly causing the early termination of eGFP expression. Consequently, fluorescence-disappeared syncytia were always observable after P40, and even predominant on cell monolayer at P50. In addition, the NGS analysis revealed the non-sense mutation with a high frequency of occurrence at P50. Even so, fluorescent syncytia were still visible at P50 (Figure 1, P50) and possibly induced by a high-fidelity mutant. If such a mutant exists in viral quasispecies at P50, its high-fidelity mechanism would be attributed to the mutated L protein, because RdRp is a primary driving force of point mutations observed in RNA virus populations (7).

Historically, a virulence-attenuated canine morbillivirus (Rockborn strain) was reported to revert back to virulent form after serial passages in dogs (27). Although there was no similar report about reversion to SRMV virulence in vivo, Abdullah et al. (28) recently demonstrated that change of a single aa was sufficient for causing SRMV to enter human cells. Therefore, screening of a high-fidelity SRMV vaccine strain would play an important role not only in immunization of animals, but also in reduction of a potential risk in viral trans-species transmission. In this study, the L sequence was found to harbor eight missense mutations during serial passaging, out of which six were identified at P40. Unexpectedly, the P50 progeny showed no SNV in the L ORF, mainly attributed to too low sequencing depths to exhibit some low-frequency mutations. We speculated that at least one high-fidelity mutant was present in viral quasispecies at P50. If purified from the population of viral progenies, this putative mutant would contribute to clarifying a molecular mechanism in viral high-fidelity replication in vitro.

**DATA AVAILABILITY STATEMENT**

The data presented in the study are deposited in the NCBI SRA repository, accession number (PRJNA736161).

**AUTHOR CONTRIBUTIONS**

FL: conceptualization and writing—original draft preparation. YZ: methodology. LL and CL: formal analysis. XW: project administration. All authors contributed to the article and approved the submitted version.

**FUNDING**

This work was supported by the National Key Research and Development Program of China (Grant Nos.: 2016YFD0500108-4, 2016YFD0500108, and 2016YFD0500100).

**ACKNOWLEDGMENTS**

We gratefully thank Yu’s group in the Shanghai Tanpu Biotechnology Co., Ltd. for the data analysis of NGS. We also thank QianQian Wang for her help in RNA extraction.

**REFERENCES**

1. Zakian A, Nouri M, Kahroba H, Mohammadian B, Mokhber-Dezfouli MR. The first report of peste des petits ruminants (PPR) in camels (Camelus dromedarius) in Iran. Trop Anim Health Prod. (2016) 48:1215–9. doi: 10.1007/s11250-016-1078-6
2. Abubakar M, Mahapatra M, Muniraju M, Arshed MJ, Khan EH, Banyard AC, et al. Serological detection of antibodies to peste des petits ruminants virus in large ruminants. Transbound Emerg Dis. (2017) 64:513–9. doi: 10.1111/tbed.12392
3. Khalafalla AI, Saeed IK, Ali YH, Abdurrahman MB, Kwiatek O, Libeau G, et al. An outbreak of peste des petits ruminants (PPR) in camels in the Sudan. Acta Trop. (2010) 116:161–5. doi: 10.1016/j.actatropica.2010.08.002
4. Mahapatra M, Parida S, Baron MD, Barrett T. Matrix protein and glycoproteins F and H of Peste-des-petits-ruminants virus function better as a homologous complex. J Gen Virol. (2006) 87:2021–9. doi: 10.1099/vir.0.81721-0
5. Barrett T, Pastoret PP, Taylor W. Rinderpest and peste des petits ruminants: virus plaques of large and small ruminants. In: Taylor WP, Roeder PL, Rweyemamu MM, editors. London: Academic Press: Elsevier (2006).
6. Parida S, Muniraju M, Mahapatra M, Muthuchelvan D, Buszkowski H, Banyard AC. Peste des petits ruminants. Vet Microbiol. (2015) 181:90–106. doi: 10.1016/j.vetmic.2015.08.009

7. Borderia AV, Rozen-Gagnon K, Vignuzzi M. Fidelity variants and RNA quasispecies. Curr Top Microbiol Immunol. (2016) 392:303–22. doi: 10.1007/82_2013_483

8. Andino R, Domingo E. Viral quasispecies. Virology. (2015) 479–480:46–51. doi: 10.1016/j.virology.2015.03.022

9. Domingo E, Sheldon J, Perales C. Viral quasispecies evolution. Microbiol Mol Biol Rev. (2012) 76:159–216. doi: 10.1128/MMBR.05023-11

10. Liu F, Wu X, Li L, Zou Y, Liu S, Wang Z. Evolutionary characteristics of morbilliviruses during serial passages in vitro: gradual attenuation of virus virulence. Comp Immunol Microbiol Infect Dis. (2016) 47:7–18. doi: 10.1016/j.cimid.2016.05.007

11. Schrag SJ, Rota PA, Bellini WJ. Spontaneous mutation rate of measles virus: direct estimation based on mutations conferring monoclonal antibody resistance. J Virol. (1999) 73:51–4. doi: 10.1128/JVI.73.1.51-54.1999

12. Zhang X, Rennick LJ, Duprex WP, Rima BK. Determination of spontaneous mutation frequencies in measles virus under nonselective conditions. J Virol. (2013) 87:2686–92. doi: 10.1128/JVI.02146-12

13. Wu X, Liu F, Li L, Zou Y, Liu S, Wang Z. Major mutation events in structural genes of peste des petits ruminants virus through serial passages in vitro: gradual attenuation of virus virulence. Comp Immunol Microbiol InfectDis (2013) 87:2686–92. doi: 10.1128/JVI.02146-12

14. Ni M, Chen C, Qian J, Xiao HX, Shi WF, Luo Y, et al. Intra-host dynamics of Ebola virus during 2014. Nat Microbiol. (2016) 1:16151. doi: 10.1038/nmicrobiol.2016.16

15. Hasing ME, Hazes B, Lee BE, Preiksaitis JK, Pang XL. A next-generation sequencing-based method to study the intra-host genetic diversity of norovirus in patients with acute and chronic infection. BMC Genomics. (2016) 17:480. doi: 10.1186/s12864-016-2831-y

16. Yang Z, Mammel M, Whitehouse CA, Ngo D, Kulkia M. Inter- and intra-host nucleotide variations in hepatitis A virus in culture and clinical samples detected by next-generation sequencing. Viruses. (2018) 10:619. doi: 10.3390/v10110619

17. Campo DS, Dimitrova Z, Yamasaki L, Skums P, Lau DT, Vaughan G, et al. Next-generation sequencing reveals large connected networks of intra-host HCV variants. BMC Genomics. (2014) 15(Suppl. 5):S4. doi: 10.1186/1471-2164-15-S5-S4

18. Liu F, Huang Y, Wang Q, Li J, Shan H. Rescue of senecavirus A to uncover mutation profiles of its progenies during 80 serial passages in vitro. Vet Microbiol. (2021) 253:108969. doi: 10.1016/j.vetmic.2020.108969

19. Li C, Wang H, Shi J, Yang D, Zhou G, Chang J, et al. Senecavirus-specific recombination assays reveal the intimate link between polymerase fidelity and RNA recombination. J Virol. (2019) 93:e00576–19. doi: 10.1128/JVI.00576-19

20. Liu F, Zhang Y, Li L, Zuo Y, Sun C, Xiaodong W, et al. Rescue of eGFP-expressing small ruminant morbillivirus for identifying susceptibilities of eight mammalian cell lines to its infection. Virus Res. (2019) 261:60–4. doi: 10.1016/j.virusres.2018.12.011

21. Crotty S, Cameron CE, Andino R. RNA virus error catastrophe: direct molecular test by using ribavirin. Proc Natl Acad Sci USA. (2001) 98:6895–901. doi: 10.1073/pnas.11085598

22. Cameron CE, Castro C. The mechanism of action of ribavirin: lethal mutagenesis of RNA virus genomes mediated by the viral RNA-dependent RNA polymerase. Curr Opin Infect Dis. (2001) 14:757–64. doi: 10.1097/00001432-200112000-00015

23. Liu F, Li L, Liu Y, Sun C, Liu C, Wu X, et al. Development of reverse genetics system for small ruminant morbillivirus: rescuing recombinant virus to express Echinococcus granulosus EG95 antigen. Virus Res. (2019) 265:50–5. doi: 10.1016/j.viruses.2018.12.008

24. Liu F, Sun C, Li L, Wang Z. Rescue of recombinant peste des petits ruminants virus for expressing Orf virus B2L protein (in Chinese). China Anim Health Inspection. (2019) 36:71–7.

25. Lu IN, Muller CP, He FQ. Applying next-generation sequencing to unravel the mutational landscape in viral quasispecies. Virus Res. (2020) 283:197963. doi: 10.1016/j.viruses.2020.197963

26. Huang SW, Hung SJ, Wang JR. Application of deep sequencing methods for inferring viral population diversity. J Virol Methods. (2019) 266:95–102. doi: 10.1016/j.jviromet.2019.01.013

27. Appel MJG. Reversion to virulence of attenuated canine distemper virus in vivo and in vitro. J Gen Virol. (1978) 41:385–93. doi: 10.1099/0022-1317-41-2-385

28. Abdullah N, Kelly JT, Graham SC, Birch J, Goncalves-Carneiro D, Mitchell T, et al. Structure-guided identification of a nonhuman morbillivirus with zoonotic potential. J Virol. (2018) 92:e01248–18. doi: 10.1128/JVI.01248-18

Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2021 Liu, Zou, Li, Liu and Wu. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.