Lineage A betacoronavirus NS2 proteins and the homologous torovirus Berne pp1a carboxy-terminal domain are phosphodiesterases that antagonize activation of RNase L

Stephen A. Goldstein  
*University of Pennsylvania*

Joshua M. Thornbrough  
*Adelphi Research Global*

Rong Zhang  
*Washington University School of Medicine*

Babal K. Jha  
*Taussig Cancer Research Institute*

Yize Li  
*University of Pennsylvania*

See next page for additional authors

Follow this and additional works at: [https://digitalcommons.wustl.edu/open_access_pubs](https://digitalcommons.wustl.edu/open_access_pubs)

Recommended Citation

Goldstein, Stephen A.; Thornbrough, Joshua M.; Zhang, Rong; Jha, Babal K.; Li, Yize; Elliott, Ruth; Quiroz-Figueroa, Katherine; Chen, Annie I.; Silverman, Robert H.; and Weiss, Susan R., "Lineage A betacoronavirus NS2 proteins and the homologous torovirus Berne pp1a carboxy-terminal domain are phosphodiesterases that antagonize activation of RNase L." *The Journal of Virology* 91,5. (2017).  
[https://digitalcommons.wustl.edu/open_access_pubs/5689](https://digitalcommons.wustl.edu/open_access_pubs/5689)
Lineage A Betacoronavirus NS2 proteins and homologous Torovirus Berne pp1a-carboxyterminal domain are phosphodiesterases that antagonize activation of RNase L

Stephen A. Goldstein1,*, Joshua M. Thornbrough1,3, Rong Zhang1,4,*, Babal K. Jha2,5, Yize Li1, Ruth Elliott1, Katherine Quiroz-Figueroa1, Annie I. Chen1, Robert H. Silverman2 and Susan R. Weiss1,#

1Department of Microbiology, Perlman School of Medicine at the University of Pennsylvania, Philadelphia, PA, USA and 2Department of Cancer Biology, Lerner Research Institute, Cleveland Clinic, Cleveland, OH, USA

Running title: Coronaviridae phosphodiesterases antagonize RNase L

Key words: Lineage A Coronavirus, Torovirus, ribonuclease L, interferon antagonism, phosphodiesterase

* These authors contributed equally to this work

Current addresses:

3 Adelphi Research Global, Doylestown, PA 18901
4 Division of Infectious Diseases, School of Medicine, Washington University in St. Louis, St. Louis, MO, 63110
5 Translational Hematology & Oncology Research; Taussig Cancer Research Institute, Cleveland Clinic, Cleveland OH, 441063

# Corresponding Author

Susan R. Weiss
Department of Microbiology
University of Pennsylvania
Perelman School of Medicine
203A Johnson Pavilion
36th Street and Hamilton Walk
Philadelphia, PA 19104-6076
Phone: 215-898-8013
FAX: 215-573-4858
Email: weisssr@upenn.edu

Word counts for the abstract: 246
Word count for text: 23,515
Abstract

Viruses in the family *Coronaviridae*, with the Nidovirus order, are etiologic agents of a range of human and animal diseases, including both mild and severe respiratory disease in humans. These viruses encode conserved replicase and structural proteins, and more diverse accessory proteins in the 3’ end of their genomes that often act as host cell antagonists. We have previously shown that 2’,5’ phosphodiesterases (PDE) encoded by the prototypical Betacoronavirus, mouse hepatitis virus (MHV), Middle East respiratory syndrome-associated coronavirus antagonize the oligoadenylate–ribonuclease L (OAS-RNase L) pathway. Here we report that additional coronavirus superfamily members including lineage A betacoronaviruses and toroviruses infecting both humans and animals encode 2’,5’ PDEs capable of antagonizing RNase L. We used a chimeric MHV system, in which exogenous PDEs were expressed from an MHV backbone lacking a functional NS2 protein (MHV<sup>Mut</sup>), its endogenous RNase L antagonist. In this system, we found that 2’,5’ PDEs encoded by human coronavirus HCoV-OC43 (OC43), an agent of the common cold, human enteric coronavirus (HECoV), equine coronavirus (ECoV), and equine torovirus-Berne (BEV) are enzymatically active, rescue replication of MHV<sup>Mut</sup> in bone marrow-derived macrophages and inhibit RNase L-mediated rRNA degradation in these cells. Additionally, PDEs encoded by OC43 and BEV rescue MHV<sup>Mut</sup> replication and restore pathogenesis in WT B6 mice. This finding expands the range of viruses known to encode antagonists of the potent OAS-RNase L antiviral pathway, highlighting its importance in a range of species, as well as the selective pressures exerted on viruses to antagonize it.
Importance

Viruses in the family Coronaviridae include important human and animal pathogens, including the recently emerged SARS-CoV and MERS-CoV. We have shown previously that two viruses within the genus Betacoronavirus murine coronavirus (MHV) and MERS-CoV, encode 2',5' phosphodiesterases (PDEs) that antagonize the OAS-RNase L pathway and report here that these proteins are furthermore conserved among additional coronavirus superfamily members including lineage A betacoronaviruses and toroviruses and suggesting they may play critical roles in pathogenesis. As there are no licensed vaccines or effective antivirals against human coronaviruses and few against those infecting animals, identifying viral proteins contributing to virulence can inform therapeutic development. Thus, this work demonstrates that a potent antagonist of host antiviral defenses is encoded by multiple and diverse viruses within Coronaviridae, presenting a possible broad-spectrum therapeutic target.
Introduction

Coronaviruses (CoV) and closely related toroviruses (ToV) are well known agents of disease in mammals, including humans. Coronaviruses and toroviruses, members of the family Coronaviridae, within the Nidovirus order, contain positive sense single stranded (ss)RNA genomes, the longest known RNA genomes ranging from 28-31kb (1). The first two thirds of their genomes encodes the replicase proteins, which include the viral RNA-dependent RNA polymerase and numerous non-structural proteins (NSPs), which are required for replication and in some cases have host immune antagonist activities (2). The structural proteins are encoded in the 3' third of the genome and consist of spike (S), small membrane protein (E), membrane (M), nucleocapsid (N) and sometimes hemagglutinin-esterase (HE). Interspersed among the structural genes are diverse genes encoding accessory proteins that are not essential for replication but are believed to be required for virulence in vivo (1).

Mouse hepatitis virus (MHV) is a lineage A Betacoronavirus and the prototypical CoV. MHV encodes the accessory protein NS2 which was previously identified as a 2-His (H) phosphoesterase (2H-PE) superfamily member (3) that we have demonstrated has 2',5'-phosphodiesterase (PDE) activity that antagonizes host interferon (IFN) signaling via antagonism of the 2',5'-oligoadenylate synthetase (OAS)-ribonuclease (RNase) L pathway (4). Upon sensing double stranded (ds)RNA, OAS synthesizes 2',5'-oligoadenylates (2-5A) which catalyze the activation of RNase L via homodimerization. RNase L subsequently cleaves host and viral ssRNA leading to termination of protein synthesis and subsequent apoptosis (5). NS2 cleaves 2-5A thus preventing the activation of RNase L. NS2 is a critical determinant of MHV strain A59 (A59) liver tropism in C57Bl/6 (B6) mice and is required for the virus to cause hepatitis. A mutant A59 (NS2H126R referred to herein as NS2Mut) expressing an inactive
phosphodiesterase is unable to antagonize the OAS-RNase L pathway in the liver of mice. Infection with this virus does not result in hepatitis and NS2<sup>Mut</sup> replication is reduced at least 10,000 fold compared to wild-type A59. However, in mice genetically deficient for RNase L (RNase L<sup>−/−</sup>) NS2<sup>Mut</sup> replicates to wild-type levels and causes hepatitis (4).

As might be expected of antagonists of a potent innate antiviral pathway, 2′,5′ PDEs are not a host evasion mechanism unique to MHV. We recently showed that the NS4b accessory protein of MERS-CoV and related bat coronaviruses, all lineage C betacoronaviruses, encode the NS4b accessory proteins with 2′,5′-PDE activity (6). Additionally, unrelated group A rotaviruses encode a PDE in the C-terminal domain of the VP3 structural protein (7). We show here that lineage A betacoronaviruses closely related to MHV, including the human respiratory HCoV-OC43 (OC43), human enteric CoV-4408 (HECoV), equine ECoV-NC99 (ECoV), and porcine hemagglutinating encephalomyelitis virus (PHEV), as well as the more distantly related equine torovirus (ETOv)-Berne (BEV) also encode NS2 homologs with predicted PDE activity. We found that these proteins do possess enzymatic 2′,5′-PDE activity that is capable of antagonizing RNase L (with the exception of the PHEV NS2) and thus countering a potent host antiviral response, suggesting that PDE mediated OAS-RNase L antagonism is an important virulence strategy for lineage A betacoronaviruses and toroviruses.

**Material and Methods**

**Cell lines and mice.** Murine fibroblast L2 (L2), murine 17 clone 1 (17Cl1) and baby hamster kidney cells expressing MHV receptor (BHK-R) were cultured as previously described (8,9). C57Bl/6 (B6) mice were originally procured from the National Cancer Institutes mouse repository, and RNase L<sup>−/−</sup> mice on a B6 genetic background were derived by Dr. Robert Silverman (10) and subsequently bred in the University of Pennsylvania animal facility.
experiments involving mice were approved by the Institutional Animal Care and Use Committee at the University of Pennsylvania. Primary bone marrow derived macrophages (BMM) were derived from marrow harvested from the hind limbs (tibia and femur) of four to six week old B6 or RNase L−/− mice as described previously (4,11). Cells were cultured in DMEM (Gibco) supplemented with 10% FBS (Hyclone) and 20% L929 cell-conditioned media for 6 days before infection.

Plasmids. NS2 genes from lineage A betacoronaviruses OC43, HECV-4408, ECoV-NC99 NC99, PHEV and pp1a-carboxyterminal domain (CTD) from the torovirus Berne were synthesized and cloned into pUC57 by BioBasic yielding pUC-OC43NS2, pUC-HECVNS2, pUC-ECoVNS2, pUC-PHEVNS2 and pUC-pp1a. The second catalytic His to Arg substitutions were made by site directed mutagenesis in all plasmids resulting in pUC-OC43NS2H129R, pUC-HECVNS2H129R, pUC-NC99NS2H129R, pUC-PHEVNS2H129R and pUC-pp1aH4516R. Select genes were subsequently subcloned into the pMal parallel-2 expression vector resulting in pMAL-OC43NS2, pMAL-OC43NS2H129R, pMAL-PHEVNS2, pMAL-PHEVNS2H129R, pMAL-pp1a and pMAL-pp1aH4516R. MHV NS2 and NS2H129R had been previously cloned into pMAL-c2 (4).

Purification of recombinant PDEs from E. coli and FRET assay. MBP-PDE fusion proteins were expressed from pMAL-plasmids in BL21 T7 expression competent E. coli (NEB, Inc., Ipswich, MA) and purified by affinity chromatography followed by ion exchange chromatography on MonoQ GL10/100 using a NaCl gradient from 0 to 1 M in 20 mM NaCl as previously described (4,12). The integrity and the purity of the purified MBP fusion proteins were determined by SDS-PAGE Coomassie Blue R250 staining. The extent of purity was similar for all of the enzymes as accessed by SDS-PAGE analysis. To assess enzymatic activity, purified proteins [10 μM MBP (420 μg/ml) as control or 1 μM OC43 (75 μg/ml); BEV (60 μg/ml) PHEV (65 μg/ml) or MHV (70μg/ml) MBP-PDE fusion proteins] in 150 μl of assay buffer (20 μM
HEPES [pH 7.2], 10 mM MgCl2, 1 mM dithiothreitol) were incubated at 30° with (2'-5')p3A3 (2-5A). After one hour, reactions were stopped by heat inactivation at 95° for 3 min followed by 30 min centrifugation at 20,000 X g (4°) and supernatants carefully removed. A fluorescent resonance energy transfer (FRET) assay was used to assess enzymatic activity by measuring the amount of uncleaved, intact 2-5A left in the reaction, as previously described (13). The abilities of recombinant enzyme to degrade 2-5A were determined by a FRET based RNase L activation assay using an authentic 2-5A (2',5'-p3A3) trimer as described earlier (4, 6, 13, 14). Assays were performed three times in triplicate using two separate enzyme preparations.

Viruses and chimeric recombinant virus construction. Wild-type MHV strain A59 and mutant NS2H126R (referred to as MHV and MHVMut in the data shown herein) were described previously (14). The chimeric viruses were constructed based on the infectious cDNA clone icMHV-A59 (8,15). The wild-type and mutant PDEs genes were PCR amplified from the pUC plasmids constructed above with primers bearing SalI and NotI restricting sites. After purification and digestion with SalI and NotI, the fragments were cloned into icMHV-A59 fragment G, with an NS2H126R mutation, as previously described (14), and confirmed by DNA sequencing. The full-length A59 genome cDNA was assembled, and the recombinant viruses were recovered in BHK-R cells as previously described (8,14,15). When virus cytopathology was observed, virus was plaque purified from the supernatant and amplified on 17CL-1 cells for use. The pairs of chimeric viruses expressing WT and mutant PDEs were named by the source of the PDE, OC43 & OC43Mut, HECoV & HECoVMut, PHEV & PHEVMut, ECoVMut & ECoV and BEV & BEVMut.

The PDE gene and flanking regions were amplified by PCR from the cloned chimeric virus genomes and the sequences verified. The primers used for sequencing were Fns4 (5'-TTGTTGTGATGAGTATGGAG) which maps 136 nucleotides upstream of the ATG start codon for the PDE and Rns4 (5'-GCGTAACCATGCATCCTCAC) which maps 139 nucleotides
downstream of the PDE ORF. The regions sequenced include the SalI and NotI restriction sites as well as the transcription regulatory sequence (TRS) for ORF4 and ORF5a.

**Chimeric MHV infections of bone marrow derived macrophages (BMM).** BMM were mock infected or infected at a multiplicity of infection (MOI) of 1 PFU/cell (in triplicate) and allowed to adsorb for 1 hour at 37°C. Cultures were washed with PBS (3 times) and fed with medium. At the times indicated, cells were lysed and analyzed for degradation of RNA (described below) or supernatants were harvested for quantification of viral titers by plaque assay on L2 cells (14).

**Immunoblotting.** L2 cells were infected with MHV or chimeric viruses (MOI=1PFU/cell). At 10 hours post infection, cells were lysed in nonidet P-40 (NP-40) buffer (1% NP-40, 2 mM EDTA, 10% glycerol, 150 mM NaCl and 50 mM Tris pH 8.0) containing protease inhibitors (Roche). Protein concentrations were measured using a DC protein assay kit (Bio-Rad). Supernatants were mixed 3:1 with 4X SDS-PAGE sample buffer. Samples were boiled, separated by 4-15% SDS-PAGE and transferred to polyvinylidene difluoride (PVDF) membranes. Blots were blocked with 5% nonfat milk and probed with the following antibodies: anti-Flag M2 mice monoclonal antibody (Agilent, 1:1000); anti MHV nucleocapsid mouse monoclonal antibody (a gift from Dr. Julian Leibowitz; 1:400) and anti GAPDH mouse monoclonal antibody (Thermo Scientific, 1:1000). Anti-mouse HRP (Santa Cruz; 1:5000) secondary antibodies were used to detect the primary antibodies. The blots were visualized using Super Signal West Dura Extended Duration Substrate (Thermo Scientific). Blots were probed sequentially with antibodies with blots being stripped between antibody treatments.

**Analysis of RNase L mediated rRNA degradation.** RNA was harvested from B6 WT BMM infected with MHV and chimeric viruses encoding WT and catalytically inactive PDEs at the indicated time points using a Qiagen RNeasy kit. RNase was denatured at 70°C for 2 min and
analyzed with an Agilent BioAnalyzer 2100 on a eukaryotic total RNA nanochip. The BioAnalyzer converts the electropherogram generated for each sample into the pseudogel as depicted in Fig 6 (4).

Replication in mice. Four week old B6 or RNase L−/− mice (5-7) were anesthetized with isoflurane (Abbott Laboratories; Chicago, IL) and inoculated intrahepatically with 2000 PFU in 50 μL of DPBS (Gibco) containing 0.75% bovine serum albumin (Sigma). Mice were euthanized with CO2, perfused with DPBS (Gibco) and livers harvested at day five post inoculation. Part of the liver was fixed for histology below and the rest was homogenized and viral titers were determined by plaque assay of liver homogenates on L2 cells (16). A piece of each liver was fixed overnight in 4% paraformaldehyde, embedded in paraffin and sectioned. Sections were stained with hematoxylin and eosin (H&E) or alternatively blocked with 10% normal donkey serum and immunostained with a 1:20 dilution of a monoclonal antibody against MHV nucleocapsid (N) protein (1:1000 dilution). Staining was developed using avidin-biotin-immunoperoxidase (Vector Laboratories).

Results

Alignment and modeling of coronavirus and torovirus NS2 proteins. To determine whether the MHV-related betacoronaviruses encode proteins with 2',5'-PDE activity we first analyzed the primary amino acid sequence of the NS2 proteins from OC43, HCoV, ECoV-NC99, PHEV and the pp1a-CTD of BEV. While the NS2 homologs are encoded within ORF2a, the PDE of BEV is encoded at the 3' end of the ORF1a and processed from the pp1a polyprotein (2). All of these proteins contain two conserved HxS/Tx motifs spaced by ~80 residues, where x is any hydrophobic residue, characteristic of 2H-phosphoesterase superfamily proteins (3,4,14) (Fig 1). Interestingly the carboxytermini of the PHEV and BEV PDEs are truncated relative to the other
NS2 proteins, similar to rotavirus VP3-CTD PDEs (14,17). We further entered the primary amino acid sequence of these proteins into Phyre² to predict their tertiary structures (Fig 2). All of these proteins scored highly for homology with the published structure of the A-kinase anchoring protein 7 (AKAP7) central domain (CD) (18), a previously identified host-encoded 2H-PE with 2′,5′-PDE (7). We have previously shown that the MHV NS2 and group A rotavirus (RVA) VP3 proteins, also structural homologs of AKAP7 CD, exhibit 2′,5′ PDE activity and can antagonize RNase L (4,7,14).

Coronavirus and torovirus putative 2′,5′ PDEs are enzymatically active and cleave 2-5A.

To determine whether the predicted Nidovirus PDEs (OC43, BEV, PHEV) are enzymatically active, the genes encoding them as well as their corresponding mutants with an Arg substitution of the second predicted catalytic His residue were expressed in Escherichia coli as maltose binding protein (MBP) fusion proteins and purified by affinity chromatography followed by ion exchange chromatography and size exclusion chromatography as described in Materials and Methods (4). Purified wild type or catalytic mutant proteins were incubated with 2-5A substrate and an indirect fluorescent resonance energy transfer (FRET) assay was used to assess activation of RNase L, in which higher RLUs represent active RNase L as described in Materials and Methods and in detail previously (13). MHV NS2 was utilized as a positive control for inhibition of RNase L (Fig 3). OC43 and BEV proteins reduced RNase L activation to a similar degree as MHV NS2, while PHEV NS2 was significantly less active. The mutant proteins containing a His → Arg mutation in the second catalytic motif did not inhibit RNase L, as expected and consistent with previously results describing MHV NS2 (4).

Coronavirus and Torovirus PDEs inhibit RNase L when are expressed from a chimeric MHV NS2 mutant backbone To investigate whether the NS2 proteins of OC43, HECoV, ECoV, PHEV, and BEV pp1a-CTD can antagonize RNase L during infection, we constructed chimeric
viruses expressing each exogenous PDE from ORF4 (encoding NS4a, 4b) of an MHV backbone (Fig 4). The MHV-A59 backbone we utilized encodes H126R substitution in NS2 (MHV\textsuperscript{Mut} referred to in literature as NS2\textsuperscript{H126R}) that abrogates its enzymatic activity and its ability to antagonize RNase L. MHV\textsuperscript{Mut} exhibits minimal replication in primary bone marrow-derived macrophages (BMM) or in vivo (4). The chimeric viruses we constructed express either the exogenous PDE protein or its catalytic mutant from the ORF4 locus of MHV, which is dispensable for MHV replication \textit{in vitro} and \textit{in vivo} (19). Each exogenous protein was constructed with a C-terminal Flag-tag to allow verification of expression from the chimeric viruses.

To assess expression of PDEs by western blot, we infected L2 cells with the chimeric viruses and harvested protein lysates at 10 hours post-infection (hpi). We probed for the exogenous PDEs using a primary antibody directed against the Flag-tag, and utilized GAPDH as a loading control (Fig 5). The OC43, HECV and ECoV PDEs were detectable by western blot at a high level of abundance, while detection of BEV pp1-CTD expression was less robust. PHEV NS2 expression from multiple viral clones as well as the swarm of uncloned recombinant virus could not be detected by western blot.

\textbf{Exogenous coronavirus and torovirus PDEs rescue replication of MHV\textsuperscript{Mut} in primary B6 BMMs through inhibition of RNase L activation.} To determine if the exogenous PDEs can antagonize RNase L in the context of infection, we infected BMMs from WT B6 and RNase L\textsuperscript{-/-} mice with MHV, MHV\textsuperscript{Mut} and the chimeric viruses and measured replication by plaque assay at 6, 9, 12, and 24 hpi. As expected, MHV\textsuperscript{Mut} is significantly attenuated in WT BMMs but replicates to equivalent titers as MHV in RNase L\textsuperscript{-/-} BMMs (Fig 6A). All of the chimeric viruses encoding WT exogenous PDEs from OC43, HECoV, ECoV and BeV, replicated to a similar extent as WT
A59 in B6 BMMs, indicating that these proteins effectively compensate for an inactive NS2<sup>H126R</sup> in MHV (Fig 4B-E). In contrast, and similarly to MHV<sup>Mut</sup>, the chimeras expressing catalytically inactive exogenous PDEs fail to replicate robustly in B6 BMMs but do replicate efficiently in RNase L<sup>-/-</sup> BMMs (Fig 6A-E). The chimeric virus encoding PHEV NS2 were not assessed for replication in BMMs due to our inability to confirm its expression (Figure 5).

To directly link antagonism of RNase L to the ability of the exogenous PDEs to rescue MHV<sup>Mut</sup> replication, we assessed rRNA degradation in infected cells by Bioanalyzer. We have previously used this assay to demonstrate that MHV NS2, but not NS2<sup>H126R</sup>, inhibits RNase L-mediated RNA degradation, and that a deficiency in RNase L obviates the requirement for NS2 in MHV replication (4). We infected B6 WT and RNase L<sup>-/-</sup> BMMs with MHV and the chimeric viruses and harvested total RNA 9 hpi. We ran the total RNA on a Bioanalyzer to visualize the integrity of rRNA during infection with MHV and the chimeric viruses. MHV and the chimeric viruses encoding exogenous PDEs encoded by MHV, OC43, HECoV, ECoV and BEV prevented rRNA degradation in B6 WT BMMs, while the corresponding catalytically inactive PDEs failed to do so (Fig 6F). This directly links the ability of the exogenous PDEs to rescue MHV<sup>Mut</sup> replication to their antagonism of RNase L activation.

OC43 NS2 and BEV pp1a-CTD restore MHV<sup>Mut</sup> replication and pathogenesis in vivo. MHV causes profound hepatitis and associated liver pathology in B6 mice, with its liver replication and pathogenesis dependent on NS2-mediated antagonism of RNase L (Fig 7) (4). To determine whether exogenous viral PDEs can rescue replication and restore pathogenesis to MHV<sup>Mut</sup>, we infected B6 and RNase L<sup>-/-</sup> mice with MHV, MHV<sup>Mut</sup> and the chimeric viruses expressing either WT or catalytic mutant PDEs from OC43 (NS2) and BEV (pp1a-CTD). Five days post-infection, at the time of peak titer, the mice were sacrificed and livers harvested for virus titration by plaque assay. In WT B6 mice chimeric viruses expressing either WT OC43...
NS2 or BEV pp1-CTD replicated robustly in the liver, similarly to MHV. In contrast, and like MHV Mut, the chimeric viruses expressing mutant OC43 NS2 (Fig 7B) or BEV pp1a-CTD (Fig 7C) are dramatically restricted, replicating only to titers below or just above the limit of detection, whereas all of the chimeric viruses replicated robustly in the livers of RNase L−/− mice (Fig 7A-C).

To assess hepatitis in these infected mice, livers sections were assessed for viral antigen and pathological changes. Like A59, chimeric viruses expressing WT OC43 NS2 or BEV pp1a-CTD caused hepatitis in B6 mice, indicated by pathologic foci in H&E stained livers, with viral antigen staining widely observed (Fig 7D,E). Chimeric viruses expressing catalytically inactive OC43 NS2 or BEV pp1a-CTD did not cause liver pathology in B6 mice and viral antigen was absent, consistent with the lack of replication (Fig 7D,E). In RNase L−/− mice, all of the chimeric viruses replicated robustly and caused pathology similar to MHV A59 (Fig 7D,E), further demonstrating that the restriction of the viruses expressing mutant PDEs in B6 mice is RNase L-mediated and that the exogenous PDEs function equivalently to MHV NS2.

**Discussion**

We have previously demonstrated 2-5A cleavage and RNase L antagonism by 2′,5′ PDEs encoded by a lineage A and a lineage C betacoronavirus (MHV and MERS-CoV respectively) and group A rotaviruses as well as by cellular AKAP7 CD (4,6,7,14). Here, we extend these findings to show that additional lineage A betacoronaviruses as well as a related torovirus family member encode 2′,5′ PDEs capable of antagonizing RNase L by cleaving 2-5A. The presence of genes encoding these proteins in multiple lineage A betacoronaviruses suggests that this gene was acquired by an ancient common ancestor of this lineage. Whether this virus was also ancestral to toroviruses and lineage C betacoronaviruses, or whether 2′,5′ PDEs were acquired by viruses in multiple independent events is unclear. The maintenance of this highly conserved protein throughout lineage A betacoronaviruses supports the idea that this protein mediates an
essential function in the diverse natural hosts of these viruses, spanning multiple mammalian families. Our finding of a homologous PDE in some groups of rotaviruses (14), a virus family, unrelated to coronaviruses is intriguing. A coronavirus recently isolated from bats was found to encode a protein likely to have originated from a bat orthoreovirus, which like rotaviruses has a dsRNA genome, suggesting the possibility of recombination between coronavirus and a dsRNA virus (20). Further support for this idea comes from a recent report of isolation of a MERS like coronavirus and a rotavirus in the feces of Korean bats (21). Additionally, the viruses encoding the PDEs we have described here infect different tissues within their hosts (1,22,23), indicating that RNase L antagonism may be required for robust replication in diverse cell types. For example, although MHV is hepatotropic, OC43 infects the upper airway, while other PDEs described here are encoded by enterotropic viruses (1,22,23).

The PDEs encoded by OC43, HECoV, ECoV and BEV antagonized RNase L and rescued replication of MHV<sub>Mut</sub> in primary WT B6 BMMs, indicating that not only are they enzymatically active 2',5' PDEs, but that they functionally compensate for an inactive MHV NS2 (Fig 3,6,7). Interestingly the BEV encoded PDE was able to antagonize RNase L and rescue virus MHV<sub>Mut</sub> replication both <i>in vitro</i> and <i>in vivo</i> despite the apparently low level of expression (Fig 5,6,7). This is not surprising as MERS NS4b PDE can rescue MHV<sub>Mut</sub> despite its very low expression level in the cytoplasm (6). PHEV NS2, is less enzymatically active than the other PDEs (Fig 3), suggesting it may be less able to antagonize RNase L. However since we could not detect expression by western blot of the PHEV PDE from a chimeric virus (Fig 5), further work will be needed to determine if it has RNase L antagonist activity in the context of an infection. Interestingly both the BEV and PHEV PDEs are truncated at the carboxytermini similar to the rotavirus PDE [Fig 3; (14)]; clearly the carboxyterminal sequences are not required for cleavage of 2-5A or RNase L antagonism as the rotavirus VP3-CTD and BEV PDEs have similar activity to MHV NS2 [Fig3,(14)]. Nevertheless, the diminished enzymatic activity of PHEV NS2 relative
to the other PDEs, suggests that while PDE may have been essential in the PHEV ancestor, it may not be required in the cells targeted by PHEV in its porcine host. However, RNase L is likely actively antiviral in other porcine tissues or stages of development, as suggested by the presence of an RNase L antagonist in protein 7 of transmissible gastroenteritis virus (24).

Although the chimeric MHVs encoding OC43-NS2 and BEV pp1a-CTD do not replicate quite as well as MHV in vivo (Fig. 6), this is unlikely due to disruption of the ORF4 gene by insertion of the exogenous PDEs as ablation of ORF4 expression within the genome of MHV strain JHM, had no effect on replication in vitro and in vivo pathogenesis and the MHV strain A59 ORF 4 is disrupted by a termination codon (25). Nevertheless, these chimeric viruses replicated robustly in vivo causing hepatitis and their respective mutants replicated to wild-type titers in the livers of RNase L−/− mice, indicating that restriction of the mutants in WT B6 mice is due to RNase L activity.

Overall, we have demonstrated that active 2′,5′ PDEs are a conserved feature of lineage A Betacoronavirus genomes, and that a homologous domain is encoded in the first open reading frame of a related nidovirus, BEV. This suggests that RNase L is a potent antiviral effector in diverse species and tissues, due to the wide host range represented by the viruses encoding these now-characterized PDEs. This thus far includes the lineage A and lineage C betacoronaviruses as well as the related toroviruses and the unrelated group A Rotaviruses (4,14). Finally since 2′,5′-PDEs are potent antagonist of host antiviral defenses encoded by multiple and diverse viruses within Coronaviridae, this class of protein may have the potential to be a broad-spectrum therapeutic target for human viruses including the OC43, a ubiquitous agent of the common cold and MERS.
Acknowledgments. Research reported in this publication was supported by National Institute of Allergy and Infectious Disease of the National Institutes of Health (NIH) under award number F32AI114143 to JMT, R21AI114920 to SRW and R01AI104887 to SRW and RHS.
References


**Figure Legends**

Fig 1. Alignment of lineage A betacoronavirus and Berne torovirus PDEs. PDEs with Genbank accession numbers are MHV NS2 (P19738.1) (26), OC43 NS2 (AAT84352.1) (27), HCoV NS2 (ACJ35484.1), ECoV NS2 (ABP87988.1) (28), PHEV NS2 (AAY68295.1) (29) and BEV (CAA36600.1) (30). Conserved catalytic HxS/Tx motifs are indicated by boxes.

Fig 2. Known and predicted structures of nidovirus PDEs. (A) Crystal structure of MHV NS2 (PDB: 4Z5V) and predicted structures of OC43 NS2 (B) and BEV pp1a-CTD (C). Predicted structures were generated using Phyre² then visualized and annotated using UCSF Chimera 1.8. Catalytic His and conserved Ser/Thr residues are indicated.

Fig 3. Assay of PDE activity of coronavirus and torovirus PDEs. Recombinant WT and mutant PDEs were incubated with 2-5A for 60 minutes and the remaining substrate was quantified using an indirect FRET based assay as described in Materials and Methods. RFU= relative fluorescence units, is proportional to 2-5A remaining. Data shown are from one representative of three independent experiments, each carried out in triplicate with separate enzyme preparations and are expressed as means ± SEM; *, P < 0.05, **, P < 0.01, ***, P < 0.001.

Fig 4. NS2 organization and construction of chimeric viruses. (A) Depiction of the NS2 protein of HCoV-OC43. Shown are the catalytic His residues at positions 49 and 129, with the His->Arg mutation shown below. (B) Genome organization of MHV with NS2 and NS4 loci indicated. Also shown are replicase open reading frames 1a and 1b, genes encoding structural proteins HE, S, E, M, N and I as well as nonstructural protein 5a. In chimeric viruses MHV NS2 residue 126 is mutated from H->R, rendering NS2 catalytically inactive (NS2^mut). The gene encoding the exogenous PDE or its catalytically inactive mutant is inserted in place of MHV NS4.
Fig 5. Expression of exogenous PDEs from chimeric viruses. L2 cells were infected with MHV or chimeric viruses and protein harvested 10 hpi and analyzed by western immunoblotting. Blots were probed with antibody against Flag to detect PDEs, anti-nucleocapsid antibody to assess chimeric viral infection and GAPDH as a protein loading control. MHV NS2 (lanes 1-2) is not Flag-tagged. Flag-tagged WT and mutant PDEs of OC43, HECV, PHEV, ECoV and BEV are detected as indicated. This blot was performed two times using proteins from independent infections with similar results.

Fig 6. Replication and activation of RNase L of chimeric viruses in bone marrow derived macrophages (BMM). (A-E) BMMs derived from WT B6 or RNase L–/– mice were infected with (A) MHV or chimeric MHV viruses expressing WT or mutant (B) OC43 NS2, (C) HECoV NS2, (D) ECoV NS2 and (E) BEV pp1a-CTD. Virus at each time point was titrated by plaque assay. Each time point is represented by three biological replicates, titrated in duplicate and variance expressed as SEM. Statistical significance was calculated by 2-way ANOVA in GraphPad Prism. **, P<0.01; ***, P<0.001. (F) Total RNA was isolated from WT B6 BMMs 9 hpi and rRNA integrity assayed using an Agilent BioAnalyzer. These data are from one of at least two independent experiments with similar results.

Fig 7. Replication and pathogenesis of chimeric viruses in vivo. (A-C) WT B6 or RNase L–/– mice (n=5-7) were infected intrahepatically with (A) MHV and MHVΔAst or chimeric viruses encoding WT or mutant (B) OC43 NS2 or (C) BEV pp1a-CTD. Five days post-infection livers were harvested and virus titrated by plaque assay. Each data point represents a single mouse liver, titrated in duplicate with variance expressed as SEM. Statistical significance determined by 1-way ANOVA in GraphPad Prism. ***, P<0.001. Liver sections from infected mice were stained with (D) H&E to identify hepatic pathology or (E) antibody to detect MHV nucleocapsid protein.