Murine AKAP7 Has a 2′,5′-Phosphodiesterase Domain That Can Complement an Inactive Murine Coronavirus ns2 Gene

Elona Gusho, a,b Rong Zhang, c Babal K. Jha, a Joshua M. Thornbrough, c Beihua Dong, a Christina Gaughan, a Ruth Elliott, c Susan R. Weiss, c Robert H. Silverman a

Department of Cancer Biology, Lerner Research Institute, Cleveland Clinic, Cleveland, Ohio, USA; Department of Biological, Geological, and Environmental Sciences, Cleveland State University, Cleveland, Ohio, USA; Department of Microbiology, Perelman School of Medicine, University of Pennsylvania, Philadelphia, Pennsylvania, USA.

E.G., R.Z., and B.K.J. contributed equally to this article.

ABSTRACT Viral 2′,5′-phosphodiesterases (2′,5′-PDEs) help disparate RNA viruses evade the antiviral activity of interferon (IFN) by degrading 2′,5′-oligoadenylate (2-5A) activators of RNase L. A kinase anchoring proteins (AKAPs) bind the regulatory subunits of protein kinase A (PKA) to localize and organize cyclic AMP (cAMP) signaling during diverse physiological processes. Among more than 43 AKAP isoforms, AKAP7 appears to be unique in its homology to viral 2′,5′-PDEs. Here we show that mouse AKAP7 rapidly degrades 2-5A with kinetics similar to that of murine coronavirus (mouse hepatitis virus [MHV]) strain A59 ns2 and human rotavirus strain WA VP3 proteins. To determine whether AKAP7 could substitute for a viral 2′,5′-PDE, we inserted AKAP7 cDNA into an MHV genome with an inactivated ns2 gene. The AKAP7 PDE domain or N-terminally truncated AKAP7 (both lacking a nuclear localization motif), but not full-length AKAP7 or a mutant, AKAP7H185R, PDE domain restored the infectivity of ns2 mutant MHV in bone marrow macrophages and in livers of infected mice. Interestingly, the AKAP7 PDE domain and N-terminally deleted AKAP7 were present in the cytoplasm (the site of MHV replication), whereas full-length AKAP7 was observed only in nuclei. We suggest the possibility that viral acquisition of the host AKAP7 PDE domain might have occurred during evolution, allowing diverse RNA viruses to antagonize the RNase L pathway.

IMPORTANCE Early virus-host interactions determine whether an infection is established, highlighting the need to understand fundamental mechanisms regulating viral pathogenesis. Recently, our laboratories reported a novel mode of regulation of the IFN antiviral response. We showed that the coronavirus MHV accessory protein ns2 antagonizes the type I IFN response, promoting viral replication and hepatitis. ns2 confers virulence by cleaving 2′,5′-oligoadenylate (2-5A) activators of RNase L in macrophages. We also reported that the rotavirus VP3 C-terminal domain (VP3-CTD) cleaves 2-5A and that it may rescue ns2 mutant MHV. Here we report that a cellular protein, AKAP7, has an analogous 2′,5′-phosphodiesterase (2′,5′-PDE) domain that is able to restore the growth of chimeric MHV expressing inactive ns2. The proviral effect requires cytoplasmic localization of the AKAP7 PDE domain. We speculate that AKAP7 is the ancestral precursor of viral proteins, such as ns2 and VP3, that degrade antiviral activity of RNase L.

Host antiviral pathways triggered by type I interferons (IFNs) are self-limiting so that after virus is eliminated, the host can restore normal cellular and tissue functions (1). Many types of viruses also prevent activation of host antiviral pathways (reviewed in reference 2). The 2′,5′-oligoadenylate (2-5A) synthetase (OAS)/RNase L system is one of the principal mediators of the IFN antiviral response (reviewed in references 3 to 6). Recently, we reported that two homologous viral proteins from unrelated viruses, coronavirus mouse hepatitis virus (MHV) strain A59 ns2 and group A rotavirus strain SA11 VP3, have 2′,5′-phosphodiesterase (2′,5′-PDE) activities that antagonize the antiviral activity of RNase L by degrading 2-5A [p(5′A2′p),5′A, where x is 1 to 3 and n is 2 or greater] (7, 8). ns2 and VP3 are eukaryotic-viral LigT-like family members that include both viral and cellular proteins of diverse origins, some of which possess cyclic nucleotide phosphodiesterase (CPD) activity (Fig. 1A) (9). LigT proteins are named for the prototypical archeo-bacterial tRNA-ligating enzyme LigT with reversible 2′-5′-RNA ligase activity (10) and are part of a larger superfamily of 2H phosphodiesterases characterized by the presence of a pair of conserved His-h-h motifs (where h is typically a hydrophobic residue) (9, 11, 12). However, while MHV ns2 has 2′,5′-PDE activity, it apparently lacks CPD activity based on its inability to cleave 2′,3′ cyclic AMP (cAMP), 3′,5′ cAMP, and ADP-ribose 1″,2″ cyclic phosphate (7). Mutation of the active site of ns2 blocked MHV replication in liver, thereby preventing hepatitis in wild-type (wt)
mice but not in RNaseL−/− mice (7). Furthermore, the group A rotavirus (strain SA11) VP3 C-terminal domain (CTD) was able to restore the replication and virulence of a chimeric ns2 mutant MHV in mice (8), showing that these two virally encoded activities are functionally equivalent. Homologous PDEs encoded by other 2a betacoronaviruses, toroviruses, and group A rotaviruses suggest that this is a general mechanism of host antagonism necessary for replication of many RNA viruses (5, 8, 9, 11; unpublished data).

A kinase anchoring proteins (AKAPs) are a family of scaffolding proteins that bind the regulatory (R) subunits of protein kinase A (PKA) to localize, coordinate, and regulate cAMP signaling during diverse processes, including cardiac excitation-contraction coupling, neuronal synaptic plasticity, sperm motility, insulin secretion, and renal homeostasis (reviewed in references 13 to 16). Muscle-specific AKAP (mAKAP) partners with a cAMP-specific phosphodiesterase, PDE4D3, which limits activation of PKA (17). Several other AKAPs also bind to different cyclic nucleotide PDEs (reviewed in reference 16). However, among more than 43 known AKAP family members (13), only long isoforms of AKAP7 (also known as AKAP15 or -18) have a central domain (CD) with two characteristic His-h-Thr/Ser-h motifs and predicted structural homology to the viral 2'-5'-PDEs MHV ns2 and rotavirus VP3 (Fig. 1A and see Fig. S1 in the supplemental material) (9, 12, 18). Therefore, we explored the possibility that instead of binding an extrinsic PDE like other AKAPs, AKAP7 might have an intrinsic PDE activity. There are four splice variants of AKAP7: (i) two short forms of 15 and 18 kDa (H9251 and H9252) that have a membrane-targeting region and the PKA binding motif but lack the nuclear localization signal (NLS) and CD and (ii) two long forms of 37 and 42 kDa (H9253 and H9254) that contain (from the N to the C terminus) an NLS, the CD, and the AKAP helix and leucine zipper that bind R subunits of PKA and also Ca2+ and Na+ channels (18). A recent report described a cre/loxP knockout in mice of AKAP7 exon 7.
from all four AKAP7 splice variants (19). AKAP7 exon 7 encodes
the C-terminally modified leucine zipper domain that binds the
RII PKA subunits, Ca^{2+} and Na^{+} channels, and the 3′ untran-
scribed region (3′-UTR). The AKAP7-deficient animals were,
however, phenotypically normal. In particular, cardiomyocytes
from AKAP7-deficient mice responded normally to adrenergic
stimulation, leading the authors to suggest that another AKAP
isofrom performs this function (19). Here we investigated the
possible role of AKAP7 in a biochemical function alternative to reg-
ulating cAMP signaling, namely, regulating 2-5A signaling to
RNase L. We determined that the AKAP7 CD is a 2′,5′-PDE that
rapidly degrades 2-5A. As a result, replication of an ns2 mutant
MHV was restored by the AKAP7 CD or by an N-terminally trun-
cated AKAP7 that retains the CD, both of which localized to the
cytosol, the site of viral replication. However, full-length AKAP7
localized to the nucleus and failed to restore replication of ns2
mutant MHV. These studies identify a novel biochemical function
of an AKAP family member and are consistent with the possibility
that the AKAP7 CD has been acquired by some viruses to evade
the antiviral activity of type I IFNs by preventing activation of
RNase L (7, 8). In addition, our findings suggest that localization
of 2-5A-degrading enzymes near sites of viral RNA synthesis may
negate the antiviral activity of RNase L.

RESULTS
Murine AKAP7 rapidly degrades 2-5A with kinetics similar to
those of MHV ns2 and rotavirus VP3. Alignment of murine
AKAP7 to its eukaryotic homologs suggests that it is a member of
an ancient family of 2H phosphoesterases that extend from plants
to humans (Fig. S2). To determine whether murine AKAP7 is a
2′,5′-PDE, cDNA encoding full-length AKAP7γ (hereinafter
“AKAP7” unless stated otherwise) was expressed in bacteria, pu-
rified, and tested for its ability to cleave the trimeric species of
2-5A, (2′,5′)-pA_3, in vitro. Wild-type (wt) AKAP7, AKAP7 mut-
tated in both conserved histidines (AKAP7H93A,H185R), and, for
comparison, MHV strain A59 ns2 and human rotavirus strain WA
VP3-CTD (Fig. 1A) were incubated with (2′,5′)-pA_3 at 37°C.
2-5A levels were measured by activation of RNase L in
vitro in comparison to a standard curve of 2-5A dilutions using a previ-
ously described fluorescence resonance energy transfer (FRET)
method (20). ns2, VP3, and AKAP7 (each at 1.5 μM) rapidly
degraded (2′,5′)-pA_3 (10 μM) such that after a 1-min incubation
at 37°C, less than 30% of the input (2′,5′)-pA_3 remained intact
(Fig. 1B). Little or no detectable 2-5A remained after 10 min of
incubation with any of the three proteins. In contrast, AKAP7
mutated in the two conserved histidine residues lacked the ability
to degrade 2-5A (Fig. 1B).

To demonstrate that AKAP7 has a 2′,5′-PDE activity that cleaves
one 5′-AMP at a time from the 2′,3′ terminus of 2-5A, we performed
incubations at a lower temperature, 22°C. As we reported previously
(7), ns2 degrades (2′,5′)-pA_3 to (2′,5′)-pA_2 and 5′-AMP, and then
the diadenylate (2′,5′)-pA_2 is degraded to 5′-AMP and 5′-ATP
(Fig. 1C). AKAP7 also removed one 5′-AMP at a time from trimeric
2-5A, and therefore it is also a bona fide 2′,5′-PDE (Fig. 1D). As
expected, active-site mutant AKAP7H93A,H185R failed to degrade
(2′,5′)-pA_3 (Fig. 1E). Quantitation of these results showed that
AKAP7 and ns2 displayed comparable kinetics (Fig. 1F).

AKAP7 degrades 2-5A in intact cells transfected with dsRNA.
To determine if AKAP7 was able to degrade 2-5A in intact cells,
cDNAs for full-length AKAP7, AKAP7 CD, and mutant
AKAP7H93A,H185R (with a C-terminal Flag epitope) were tran-
siently expressed in the human ovarian carcinoma cell line Hey1B
(21). Hey1B cells were selected for these experiments because en-
dogenous AKAP7 was undetectable by Western blotting with a
rabbit polyclonal antibody against an AKAP7 CD peptide
(Fig. 1G). Ectopic expression of the different AKAP7 proteins was
confirmed by immunoblotting (Fig. 1G). At 20 h posttransfection
with either an empty vector or the AKAP7 cDNAs, cells were
transfected for an additional 3 h with the synthetic double-
stranded RNA (dsRNA) poly(rI)-poly(rC) (pIC), a potent activ-
ator of OAS (22). 2-5A was undetectable in control cells trans-
fected with the vector alone as determined by FRET (the lower
limit of detection was about 15 fmol/10⁶ cells) (Fig. 1H). However,
pIC caused high levels of 2-5A (38 pmol/10⁶ cells) to accumulate
in the vector control cells. In contrast, expression of the AKAP7
CD reduced pIC induction of 2-5A by almost 200-fold (to
0.2 pmol/10⁶ cells). The full-length AKAP7 reduced pIC-induced levels of 2-5A by about 12-fold (to 3 pmol/10⁶ cells), while full-
length mutant AKAP7H93A,H185R failed to deplete pIC-induced levels of 2-5A (26 pmol/10⁶ cells remaining). These findings dem-
onstrate that AKAP7 (full length and CD) effectively degraded
2-5A in intact pIC-transfected cells.

Viral expression of AKAP7 in BMM. To determine if AKAP7
could functionally replace ns2, we further exploited an MHV-A59
reverse-genetics system (7, 8, 23). We hypothesized that the ex-
pression of AKAP7 from a chimeric ns2 mutant MHV would re-
store viral replication in B6 bone marrow-derived macrophages
(BMM). Four MHV-AKAP7 chimeric viruses, all based on the
ns2H126R mutant MHV, were constructed (Fig. 2A) by inserting
coding sequences for full-length AKAP7 (ns2H126R-AKAP7),
AKAP7 with an N-terminal domain (NTD) truncation (ns2H126R-
AKAP7NTD), the AKAP7 CD (ns2H126R-AKAP7CD), or a mutant
AKAP7 CD (ns2H126R-AKAP7CD H185R) into the MHV nonessen-
tial ns4 gene (Fig. 2A). BMM (Rnasel-/−) were infected with each
chimeric virus, as well as with control wt A59 and ns2H126R,
for 10 h at a multiplicity of infection (MOI) of 1. All viruses expressed
similar levels of ns2 and AKAP7 proteins of the expected sizes in
infected BMM as determined by Western blotting (Fig. 2B).

Intracellular localization of wt and mutant AKAP7 proteins.
Full-length AKAP7 has an NLS sequence at its N terminus which was
deleted from the other AKAP7 constructs (Fig. 1A and 2A). We
compared the subcellular localizations of the AKAP7 proteins
expressed by each of the chimeric viruses. Indirect immunofluo-
rescence analysis with antibody against an AKAP7 CD peptide
was carried out in 17CL-1 fibroblasts and BMM, neither of which ex-
pressed a detectable level of endogenous AKAP7. Full-length
AKAP7 was located in the nuclei of both 17C1-1 and B6 BMM, as
was previously reported (Fig. 3A and B) (24). The N-terminal
deletion (NTD) including the NLS caused the truncated AKAP7
protein to localize exclusively to the cytoplasm of 17C1-1 cells but
to both the nuclei and the cytoplasm of BMM (Fig. 3A and B and
data not shown). The CD of AKAP7 and its mutant (also missing
the NLS) localized to the cytoplasm of 17C1-1 cells but, like
AKAP7NTD-, localized to both the cytoplasm and the nuclei of
BMM. Likewise, the ns2 protein was clearly cytoplasmic in 17C1-1
cells but present in both the nuclei and the cytoplasm of BMM. To
confirm these findings, human 293T cells were stably transfected
with cDNA expressing full-length AKAP7, the AKAP7 CD, or, as
a control, the empty vector. Western blots showed that the 293T
cells also failed to express detectable levels of endogenous AKAP7,
whereas both AKAP7 and AKAP7 CD were clearly present in the transfected cells (Fig. 3C). As expected, full-length AKAP7 localized to the nuclei, whereas AKAP7 CD, which lacks the NLS, localized to the cytoplasm (Fig. 3D). These studies show that expressed full-length AKAP7 is nuclear but that deletion of its NLS causes some or all of the truncated protein to localize to the cytoplasm, depending on the cell type.

**AKAP7 polypeptides that localize to the cytoplasm restore the replicative capacity of ns2 mutant MHV in BMM of wt B6 mice.** To determine the effect of AKAP7 on viral replication, BMM from wt B6 or Rnasel−/− B6 mice were infected with AKAP7 chimeric viruses or with control wt A59 and mutant ns2H126R. As expected, the replication of ns2H126R was severely impaired (by 3 log10 units) compared to that of wt A59 in B6 BMM (Fig. 4A), consistent with the results of our previous studies (7, 8, 25). Intriguingly, chimeric ns2H126R-AKAP7 was unable to rescue replication in B6 BMM, demonstrating that expression of full-length AKAP7 is nuclear but that deletion of its NLS causes some or all of the truncated protein to localize to the cytoplasm, depending on the cell type.

**Expression of the N-terminally deleted AKAP7 or the CD of AKAP7 inhibits RNase L-mediated rRNA degradation induced by the ns2 mutant virus in B6 BMM.** To further investigate whether the restoration of ns2 mutant virus replication by AKAP7 CD or N-terminally deleted AKAP7 expression is due to antagonism...
of RNase L activity, we analyzed the integrity of rRNA in BMM infected with each recombinant virus. RNase L cleaves rRNA in intact ribosomes, resulting in a characteristic set of discrete rRNA cleavage products (26, 27). BMM infected with each virus were lysed at 10 h postinfection, and RNA was extracted and analyzed with an RNA chip (see Materials and Methods) (28). The chimeric viruses expressing either the AKAP7 N-terminally truncated protein or the CD of AKAP7 prevented RNase L activity, whereas full-length AKAP7, which localizes to the nucleus, or mutant AKAP7 CD does not.

Expression of the AKAP7 CD enhances replication of ns2 mutant virus in liver. We have previously shown that the ns2 mutant is highly attenuated for replication and pathogenesis in the livers of B6 mice, but it replicated and induced hepatitis to an extent similar to that of wt A59 in Rnasel−/− mice (7, 12, 25). To determine if expression of AKAP7 is able to compensate for an inactive ns2 protein and confer liver replication in vivo, B6 and Rnasel−/− mice were infected intrahepatically with viruses expressing AKAP7 CD or its mutant as well as A59 and ns2H126R.
The virus titers in liver were determined at day 5 postinfection, the peak day for viral replication in this organ. As expected, ns2H126R virus replicated minimally in B6 mice but recovered to wt A59 virus replication levels in Rnasel−/− mice (Fig. 4D). The chimeric ns2H126R-AKAP7CD virus replicated in B6 mice to a titer of 10^4 PFU/g tissue, while the isogenic AKAP7 mutant ns2H126R-AKAP7CD H185R virus failed to replicate above the level of detection (P < 0.001) (Fig. 4E). Both of the chimeric AKAP7 viruses replicated equally in Rnasel−/− mice to a level similar to that of the ns2H126R-AKAP7CD virus in B6 mice. These results suggested that expression of the active AKAP7 CD promotes the replication of mutant ns2 virus in vivo as a result of its 2′,5′-PDE activity, which cleaves 2-5A and thereby prevents RNase L activation in the cytoplasm.

**DISCUSSION**

Control of viral infections by regulation of 2′,5′-A turnover. Our findings establish the AKAP7 CD as a 2′,5′-PDE that is able to substitute for a viral enzyme, MHV ns2, with the same activity. 2′,5′-PDEs that cleave 2-5A to ATP and AMP and phosphatases that remove the 5′-terminal phosphates on 2-5A prevent perpetual activation of RNase L after the viral infection is cleared and thereby limit RNA damage to the host (29, 30) (reviewed in reference 5). In vitro, enzymes with 5′-phosphatase activity, such as alkaline phosphatase, can remove the 5′-triphosphate moiety of 2′,5′-A, thus eliminating, or greatly reducing, the ability of the core 2′,5′-oligoadenylate to activate RNase L (31). In addition, porcine coronavirus transmissible gastroenteritis virus (TGEV) gene 7 protein has been reported to dephosphorylate 2′,5′-A through its interactions with protein phosphatase PP1 (32). Two mammalian PDEs, PDE12 (2′,5′-PDE) (33–35) in the exonuclease-endonuclease-phosphatase family of deadenylases and ectonucleotide pyrophosphatase/phosphodiesterase 1 (ENPP1) (36), have been shown to degrade 2′,5′-A. PDE12 is a mitochondrial protein with both 2′,5′- and 3′,5′-PDE activities that removes poly(A) tails from mitochondrial mRNAs (34, 35). ENPP1 has a catalytic domain that is extracellular and is therefore essentially an extracellular enzyme; in addition to degrading 2′,5′-RNA, DNA, and cAMP (36). In contrast to PDE12 and ENPP1, which have mitochondrial and extracellular locations, respectively, OAS proteins localize to either the cytoplasm or nuclei, also sites of virus replication (37, 38). Our findings here demonstrate that AKAP7 is an additional host enzyme with 2′,5′-PDE activity. Among the host 2′,5′-PDEs described to date, only AKAP7 is a 2H-phosphoesterase family member with homology to the viral enzymes of the same class.

A prior study with a commercial AKAP7 antibody (from Pro-
teintech) showed widespread expression of AKAP7 in different mouse organs, including heart, brain, skeletal muscle, kidney, and lung (19). Using the same antibody, we were able to detect low levels of AKAP7 in Hey1b cells and BMM. However, AKAP7 mRNA is not induced by wt A59 or ns2 mutant infection (unpublished data). Thus, AKAP7 may be functional at a low expression level, may be expressed at higher levels in some tissues (presently under investigation), and/or may be induced by cytokines or other soluble mediators not induced by MHV.

Cytoplasmic expression of the AKAP7 PDE rescues ns2 mutant MHV. AKAP7 is one of a relatively few members of the AKAP family that localizes to the nucleus (24, 39). Although expressed full-length AKAP7 traffics to nuclei, the AKAP7 central PDE domain typically localizes to the cytoplasm, or to both the cytoplasm and nuclei, after the N-terminal NLS is deleted (Fig. 3) (24). 2-5A produced in the cytoplasm during viral infections might be expected to transit nuclear pores, leading to its degradation by endogenous nuclear AKAP7. However, expression of full-length AKAP7 failed to rescue ns2 mutant MHV or to prevent RNase L activation in the infected wt BMM. In contrast, the AKAP7 CD rapidly degraded 2-5A, preventing activation of RNase L and restoring the ability of an ns2 mutant MHV to replicate in vitro and in vivo. MHV RNA replication occurs completely in the cytoplasm of transfected cell lines (40), and the local requirement for degradation of 2-5A may not be so surprising. Nilsen and Baglioni (41) proposed, based in part on their experiments with encephalomyocarditis virus, that microdomains (localized accumulations) of 2-5A occur at the sites of viral double-stranded replication intermediates (RI), where OAS binds and is activated, causing localized activation of RNase L. Thus, 2′,5′-PDE activity may be required in the same subcellular compartment as viral RNA replication to effectively antagonize RNase L activation, as measured by virus rescue as well as protection of rRNA. These findings raise the possibility of an additional, yet-to-be-identified isoform of AKAP7 that retains the CD but localizes to the cytoplasm. Alternatively, AKAP7 may be relevant to viruses that replicate in nuclei but not to viruses, such as coronaviruses or rotaviruses, that replicate in the cytoplasm. It remains to be determined, however, whether the endogenous, nuclear AKAP7 has any role in viral infections. Alternatively, it is also possible that AKAP7 functions to degrade nuclear 2′-5A or 2′-5A-like molecules that might be produced during nonviral cellular responses to stress (42). However, full-length AKAP7 was able to degrade 2-5A in cells transfected with pIC, although less effectively than the AKAP7 CD, perhaps due to some leakage of the overexpressed AKAP7 into the cytoplasm (Fig. 1H). Moreover, the intracellular localizations of transfected pIC and the MHV RI are likely to be different. In addition, posttranslational modifications, such as phosphorylation, may cause AKAP7 to relocate to different intracellular sites, as suggested for the AKAP species Chd8 (39).

The titers of ns2 mutant chimeric viruses with AKAP7 genes inserted in the place of MHV gene 4 are all significantly lower than those with the natural gene 4 sequences (Fig. 4D and E). This is similar to our findings with VP3-CTD/MHV chimeric viruses, which reached liver titers similar to those of the AKAP7-CD chimeras (8). The reduction in viral titers is most likely due to attenuation associated with disturbing the genome in the region of gene 4, which is not required for viral replication or virulence (43). This explanation is supported by our finding that a virus expressing wt ns2 and mutant VP3-CTD from the gene 4 position is as attenuated as virus expressing mutant ns2 and wt VP3-CTD from the gene 4 position (data not shown). Because the replication of these chimeric viruses in the liver is so much less robust than that of wt A59, it was not possible to determine whether expression of foreign PDEs fully confers hepatitis on the ns2<sup>1126R</sup> mutant.

Coronavirus replication is reported to occur only in the cytoplasm of infected cells (40) and to involve the rearrangement of cellular membranes into double-membrane vesicles and convoluted membranes, the sites of viral RNA replication (44, 45). As expected, in infected murine fibroblasts, coronavirus proteins are localized in the cytoplasm, as shown in Fig. 3A. Most, if not all, studies of localization of MHV-encoded proteins or membrane rearrangement have been carried out with transformed cell lines, and there is little information on infection of primary cells such as BMM. Interestingly, full-length AKAP7 retains its exclusively nuclear localization in BMM, whereas the N-terminally deleted AKAP7 and AKAP7 CD as well as ns2 were present in both the cytoplasm and the nucleus (Fig. 3B). Future studies will be directed at comparing the subcellular localization of proteins and replication complexes in MHV-infected primary cells to those in transformed cell lines. Nevertheless, rescue of the ns2 mutant phenotype occurs only when at least some of the AKAP PDE is localized to the cytoplasm.

Relationship between viral and cellular eukaryotic-viral LigT-like members of the 2H phosphoesterase superfamily. Degradation of 2-5A appears to be a general strategy of many RNA viruses for preventing activation of RNase L, which would otherwise block viral replication. Group 2a betacoronaviruses, which have plus-strand RNA genomes, and the group A rotaviruses, with segmented dsRNA genomes, are unrelated RNA viruses with different replication strategies, yet both encode related 2′,5′-PDEs, ns2 and VP3, respectively (7, 8). Here we present the first evidence that a viral pathogen for humans, the rotavirus strain WA (46), also encodes a functional 2′,5′-PDE (Fig. 1B). In addition, many related viruses encode predicted or confirmed 2′,5′-PDEs, including additional 2a betacoronaviruses, human OC43 and HEC4408, bovine coronavirus, porcine hemagglutinating virus, the torovirus and coronavirus superfamily member equine torovirus (Berne), and group A, B, and G rotaviruses (reviewed in reference 5). While the evolutionary origins of these viral 2′,5′-PDEs are unknown, we suggest that the host AKAP7 2′,5′-PDE domain coding sequence might have been captured through RNA recombination during viral infections in the distant past. Alternatively, the viral PDEs may have evolved from other vertebrate 2H phosphoesterases (9). The phylogenetic relationship of different species of AKAP7, VP3 and ns2, based on protein sequence data was explored by means of fast minimum evolution (see Fig. S3 in the supplemental material). While this analysis shows a close relationship among different species of AKAP7, VP3, pp1a, and ns2, it does not reveal whether this relationship is the result of divergent or convergent evolution (Fig. S3). Coronavirus mRNAs are transcribed from their genomes by a discontinuous process believed to involve switching of template by the viral replicase complex. During this process, there is a high rate of homologous recombination, reportedly up to 25%, observed during in vitro replication and in vivo (reviewed in reference 47); this high-frequency template switching may result in low-frequency copying of host mRNA. Indeed, sequences homologous to the 5′-end of major histocompatibility complex (MHC) class I coding regions are found on the 5′-end of the MHV HE gene, adjacent to the 3′-end of the ns2 open readingframe.
reading frame, and it was previously speculated that this was a result of homologous recombination between the MHV genome and host RNA (48). We suggest that AKAP7 PDE sequences have been acquired by a similar process and subsequently evolved into ns2. In support of this idea, an alignment of the amino acid sequences of different isoforms of AKAP7, ns2 and VP3, show extensive homology that extends beyond the His-h-Thr/Ser-h motifs (Fig. S1). Subsequent to randomly acquiring the AKAP7 CD coding sequence, the virus would have a selective advantage resulting in retention of the gene.

MATERIALS AND METHODS

Cell lines and mice. Murine 17C1-1, L2 fibroblast, and BHK MHV receptor (MHVR) cells were cultured as described previously (12, 23). Human 293T (ATCC) and human Hey1b (21) cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM) and RPMI 1640, respectively, both with 10% fetal bovine serum (FBS). Primary BMM were generated from the hind limbs of C57Bl/6 (B6) or ICR 5- to 7-week-old mice and cultured as described previously (49). C57Bl/6 (B6) mice were purchased from the Jackson Laboratory. ICR 5- to 7-week-old mice (B6, 10 generations of backcrossing) (50, 51) were bred in the University of Pennsylvania animal facility under an approved IACUC protocol.

Antibodies. The following antibodies were used: mouse anti-nS2 monoclonal antibody (provided by Stuart Siddell, University of Bristol, United Kingdom), mouse monoclonal anti-Flag epitope (M2; Sigma); sheep anti-mouse IgG and horseradish peroxidase (HRP)-linked whole antibody (GE Healthcare); horse anti-mouse IgG HRP-linked antibody, goat anti-rabbit IgG HRP-linked antibody (Cell Signaling); Alexa Fluor 488 goat anti-mouse IgG, Alexa Fluor 594 goat anti-rabbit IgG secondary antibodies, and Alexa Fluor 488 goat anti-rabbit IgG (Invitrogen); mouse anti-glyceraldehyde 3-phosphate dehydrogenase (GAPDH) (U.S. Biolog-
phosphate-buffered saline (PBS), lysed in buffer (50 mM Tris-HCl, pH 7.2, 0.15 M NaCl, 1% NP-40, 200 μM sodium orthovanadate, 2 mM EDTA, 5 mM MgCl₂, 5 mM DTT) and heated to 95°C for 7 min. Lysates were centrifuged for 10 min at 14,000 × g at room temperature, and supernatants were passed through Microcon centrifugal filters with a molecular mass cutoff of 3 kDa (Millipore Corporation) for 45 min at 11,000 × g. Levels of 2-5A were determined by RNase L-based FRET assays in comparison to a standard curve of authentic (2'-5')pApA, as we described previously (20).

**Viral growth kinetics.** Viral growth curves were determined by infecting B6 or Rnasel-/- MDM with each virus at an MOI of 1 PFU/cell. After a 1-h incubation, the cells were washed with PBS and cultured with DMEM supplemented with 10% FBS. Culture supernatants were collected at 6, 9, 12, and 24 h postinfection, and virus titers were determined by plaque assays on L2 cells (56).

**RNA chip analysis.** For analysis of RNA integrity, total cellular RNA was isolated at 10 h postinfection with the RNaseasy kit (QIAGEN). RNA was quantified with a NanoDrop analyzer, and equal quantities of RNA were resolved on RNA chips using an Agilent 2100 bioanalyzer (28).

**Immunofluorescence assays.** 17Cl-1 cells or B6 MDM were infected with each virus at an MOI of 1. At 9 h postinfection, the cells were washed with PBS and lysed in buffer (50 mM Tris-HCl, 15 mM EDTA, 5 mM MgCl₂, 5 mM DTT) and heated to 95°C for 7 min. Lysates previously (20).

**Western blotting.** Proteins in cell lysates were separated in 12.5% SDS-polyacrylamide gels and transferred to polyvinylidene difluoride (PVDF) membranes. Membranes were blocked and probed with the antibodies described above and developed using Amersham ECL Western blotting detection reagent (GE Healthcare) and X-ray film (Fig. 1G and 3C) or using Western Lightning Plus-ECL enhanced-chemiluminescence substrate (PerkinElmer), and proteins were detected under an Intelligent dark box II (FujiFilm) (Fig. 2B). As controls for loading and transfer, the blots were probed with anti-glyceraldehyde-3-phosphate dehydrogenase (GAPDH) or anti-β-actin.

**Animal experiments.** Four-week-old B6 or Rnasel-/- mice were anesthetized with isoflurane (IsoFlo; Abbott Laboratories) and inoculated intrahepatically with each virus (2,000 PFU/mouse) in 50 μl of PBS containing 0.75% BSA. At 5 days postinfection, the mice were sacrificed and perfused with PBS. The livers were removed, homogenized, and titrated by plaque assays on L2 cells as previously described (57). All mouse experiments were reviewed and approved by the University of Pennsylvania IACUC.

**Statistical analysis.** A two-tailed t test was performed to determine statistical significance, and the P values are shown in Fig. 4. Data were analyzed with GraphPad Prism software (GraphPad Software, Inc., CA).

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at http://mbio.asm.org/lookup/suppl/doi:10.1128/mBio.01312-14/-/DCSupplemental.

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