Selective Packaging in Murine Coronavirus Promotes Virulence by Limiting Type I Interferon Responses

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ABSTRACT Selective packaging is a mechanism used by multiple virus families to specifically incorporate genomic RNA (gRNA) into virions and exclude other types of RNA. Lineage A betacoronaviruses incorporate a 95-bp stem-loop structure, the packaging signal (PS), into the nsp15 locus of ORF1b that is both necessary and sufficient for the packaging of RNAs. However, unlike other viral PSs, where mutations generally resulted in viral replication defects, mutation of the coronavirus (CoV) PS results in large increases in subgenomic RNA packaging with minimal effects on gRNA packaging in vitro and on viral titers. Here, we show that selective packaging is also required for viral evasion of the innate immune response and optimal pathogenicity. We engineered two distinct PS mutants in two different strains of murine hepatitis virus (MHV) that packaged increased levels of subgenomic RNAs, negative-sense genomic RNA, and even cellular RNAs. All PS mutant viruses replicated normally in vitro but caused dramatically reduced lethality and weight loss in vivo. PS mutant virus infection of bone marrow-derived macrophages resulted in increased interferon (IFN) production, indicating that the innate immune system limited the replication and/or pathogenesis of PS mutant viruses in vivo. PS mutant viruses remained attenuated in MAVS−/− and Toll-like receptor 7-knockout (TLR7−/−) mice, two well-known RNA sensors for CoVs, but virulence was restored in interferon alpha/beta receptor-knockout (IFNAR−/−) mice or in MAVS−/− mice treated with IFNAR-blocking antibodies. Together, these data indicate that coronaviruses promote virulence by utilizing selective packaging to avoid innate immune detection.

IMPORTANCE Coronaviruses (CoVs) produce many types of RNA molecules during their replication cycle, including both positive- and negative-sense genomic and subgenomic RNAs. Despite this, coronaviruses selectively package only positive-sense genomic RNA into their virions. Why CoVs selectively package their genomic RNA is not clear, as disruption of the packaging signal in MHV, which leads to loss of selective packaging, does not affect genomic RNA packaging or virus replication in cultured cells. This contrasts with other viruses, where disruption of selective packaging generally leads to altered replication. Here, we demonstrate that in the absence of selective packaging, the virulence of MHV was significantly reduced. Importantly, virulence was restored in the absence of interferon signaling, indicating that selective packaging is a mechanism used by CoVs to escape innate immune detection.
Coronaviruses (CoVs), like all positive-single-stranded RNA (+ssRNA) viruses, replicate in the cytoplasm of the host cell on restructured membranes (1, 2). In the case of coronaviruses, replication/transcription complexes (RTCs) assemble on endoplasmic reticulum (ER) membranes. The RTC, a multiprotein complex, replicates genomic RNA (gRNA) and transcribes subgenomic RNA (sgRNA) during viral infection. During replication, different viral RNAs are replicated, transcribed, translated, or assembled into virions. In RNA virus infections, replication and assembly are often coordinated by cis-acting RNA elements, which may undergo intramolecular RNA interactions to form secondary structures important for recognition. Concordantly, during coronavirus infection, gRNA and sgRNAs must be distinguishable for the purposes of replication, transcription, translation, and assembly.

CoVs are very adept at packaging only gRNA into virions, while sgRNAs are almost completely excluded from virions, despite having identical 5' and 3' ends as the gRNA. In lineage A betacoronaviruses (β-CoVs), such as murine hepatitis virus (MHV), the selective packaging of gRNA is dependent on the packaging signal (PS) and its interaction with structural proteins. Early studies of the MHV PS identified a 190-bp region within the nsp15 locus as the PS (3). In both infection and virus-like particle systems, the 190-bp PS was sufficient for packaging of nonviral RNA (4). Selective packaging is intimately tied to interactions with the structural proteins N and M. The selective packaging of nonviral RNA was correlated with M protein binding (5). Furthermore, N protein binds to RNA indiscriminately and was not required for packaging of nonviral RNA. However, when the N protein C-terminal domain (CTD) from MHV was swapped with sudden acute respiratory syndrome-associated CoV (SARS-CoV), significant increases in packaged sgRNAs were observed (6). These two studies propose two different models for selective packaging with M, or alternatively N, protein playing the dominant role in conferring packaging selectivity.

A subsequent report on the PS identified a 95-bp stem-loop structure within the previously described 190-bp region (7). Structural study of the PS from MHV, human coronavirus HKU1 (HCoV-HKU1), bovine CoV, and HCoV-OC43 revealed several conserved elements within the 95-bp stem-loop structure, including 2-nucleotide (nt) bulges, AGC/GUAUU motifs, and a pentaloop (Fig. 1A) (7). More recent studies utilizing 20 synonymous mutations predicted to abolish the PS stem-loop structure demonstrated that the secondary structure of the PS was necessary for selective packaging during MHV infection (8). In this instance, significantly higher levels of sgRNA were packaged into PS mutants than in wild-type MHV. Remarkably, this increase in sgRNA did not accompany equivalent decreases in packaging of gRNA or in viral titers (8). Thus, these PS mutants had no defect in virus production in tissue culture cells compared to wild-type MHV (8). This is in contrast to the role of the alphavirus packaging signal, where PS mutants also lead to increased sgRNA packaging, but at the expense of gRNA packaging and viral fitness in cultured cells (9). Retrovirus packaging signal mutants also lead to greatly decreased genomic RNA packaging with a concomitant increase in cellular mRNA packaging (10).

Since PS mutations in non-CoVs result in large decreases in gRNA incorporation into virions and diminished replication, infection with lineage A betacoronaviruses provides a unique opportunity to study the effect of selective viral packaging in vivo. Here, using two different PS mutants, we demonstrate that disrupting the selective packaging of MHV resulted in highly attenuated viruses in vivo, and this was true for A59 and JHMV, two different strains of MHV. Furthermore, the PS was important for repressing the type I interferon (IFN-I) response during MHV infection of bone marrow-derived macrophages (BMMs). Finally, IFN-I signaling was essential for the attenuation of selective packaging mutants as the virulence of P/S mutants was restored in interferon alpha/beta receptor-knockout (IFNAR−/−) mice or mice treated with IFNAR-blocking antibody.
This study establishes the PS as a novel virulence factor for MHV, and likely other coronaviruses.

RESULTS

MHV packaging signal mutants are attenuated in vivo. Previously, we inserted the RNA sequence encoding a hemagglutinin (HA) tag within the PS of the nsp15 locus (shown in Fig. 1A and B). This virus, termed rA59Nsp15-HA, replicated like wild-type virus but had increased levels of packaged sgRNA, indicating that selective packaging of gRNA was disrupted (11). Based on these observations, we next investigated the effect of selective packaging on virulence. B6 male mice were infected with rA59Nsp15-HA by intracranial injection and monitored for morbidity and mortality for 12 days postinfection (dpi). rA59Nsp15-HA-infected mice had decreased weight loss and increased survival.

**FIG 1** rJHMV packaging mutants are attenuated in vivo. (A) The RNA secondary structure of the MHV packaging signal (7). (B) Mfold predicted RNA secondary structure of rA59Nsp15-HA. HA epitope coding sequence and complement are boldfaced and underlined. (C) Mfold predicted RNA secondary structure of MHV PS. Synonymous mutations are boldfaced and underlined. (D to F) Five- to 8-week-old male B6 mice were infected with $3 \times 10^4$ PFU of rJHMV$_{nsp15-HA}$ or rJHMV$_{WT}$ by intranasal inoculation. Infected mice were monitored for survival (D) and weight loss (E) for 14 dpi (rJHMV$_{WT}$, $n = 10$; rJHMV$_{nsp15-HA}$, $n = 15$). (F) Infected brains were harvested from mice at the indicated day postinfection. Brains were homogenized in PBS, and titers of infectious virus were determined by plaque assay on HeLa-MHV R cells. (G to I) Five- to 8-week-old male B6 mice were infected with $3 \times 10^4$ PFU of either rJHMV$_{PS}$ or rJHMV$_{WT}$ by intranasal inoculation. Infected mice were monitored for survival (G) and weight loss (H) for 14 dpi (rJHMV$_{WT}$, $n = 15$; rJHMV$_{PS}$, $n = 20$). (I) Infected brains were harvested from mice at the indicated day postinfection. Brains were homogenized in PBS, and titers of infectious virus were determined by plaque assay on HeLa-MHV R cells. The dashed line in panels F and I represents the limit of detection for the plaque assay. *, $P < 0.05$; **, $P < 0.01$; ***, $P < 0.001$ by Mann-Whitney test.
compared to wild-type rA59 (rA59WT) (see Fig. S1A and B in the supplemental material). To test whether this phenotype was strain specific, we introduced the HA sequence into nsp15 of the rJHMV strain of MHV (rJHMV-Nsp15-HA), using a previously described full-length pBAC-based reverse genetics system (12). rJHMV-Nsp15-HA had growth kinetics in vitro equivalent to that of rJHMVWT (Fig. S2A), as previously described for rA59Nsp15-HA-infected cells (11). Mice infected with rJHMV-Nsp15-HA had substantially decreased weight loss and increased survival (Fig. 1D and E). While these data suggest that selective packaging is important for MHV virulence, we cannot rule out subtle effects of the HA tag on the function of nsp15 endoribonuclease, a known IFN-I antagonist. Further, addition of the HA sequence resulted in a 31-bp double-stranded RNA (dsRNA) stem, which could conceivably induce an innate immune response (13, 14).

To separate the roles of nsp15 protein and of the PS in virulence, we constructed a second set of viruses with packaging defects in the context of both rA59 and rJHMV (termed PS-), as previously described (8). These mutants have synonymous mutations throughout the PS, which are predicted to alter the stem-loop structure of the PS without changing the amino acid sequence of nsp15 (Fig. 1C). The growth kinetics of two independent rJHMV-PS- clones (3.2 and 4.2) was nearly identical to that of rJHMVWT in vitro (Fig. S2B). These data are consistent with previous results using rA59-PS- and indicate that the PS has few, if any, effects on replication in vitro (8). After confirming the fitness of rJHMV-PS- in vitro, we next assessed the virulence of rJHMV-PS- in infected mice. B6 mice were infected intranasally with either rJHMVWT or rJHMV-PS- and monitored for morbidity and mortality for 14 dpi. Intranasal infection of rJHMVWT resulted in nearly 100% mortality (Fig. 1G) and substantial weight loss (Fig. 1H). Mice infected with rJHMV-PS- had significantly increased survival (Fig. 1G) and reduced weight loss (Fig. 1H). In agreement with results using rJHMV, mice infected with rA59-PS- also had decreased weight loss and increased survival compared to mice infected with rA59WT (Fig. S1C and D).

We next measured the viral load of rJHMVWT, rJHMV-PS-, and rJHMV-Nsp15-HA in the brains of infected mice. Both rJHMVWT and rJHMV-Nsp15-HA invaded the central nervous system (CNS) within 3 dpi and replicated to similar levels up to 5 dpi (Fig. 1F). By 7 dpi, titers in rJHMV-Nsp15-HA-infected mice were decreased in the CNS compared to those in mice infected with rJHMVWT. rJHMV-Nsp15-HA was cleared by 9 dpi. rJHMV-PS- also invaded the CNS, albeit with a delay, and reached similar peak titers as rJHMV-Nsp15-HA but not rJHMVWT (Fig. 1I). rJHMV-PS- titers decreased on 9 dpi and were cleared by 11 dpi (Fig. 1I). Thus, both PS mutant viruses did not reach the same peak viral titers as, and were cleared more rapidly than, wild-type JHMV. These data, utilizing two distinct PS mutant viruses, indicate that selective packaging of viral genomic RNA is essential for optimal virulence in multiple models of MHV infection.

rA59-PS- and rA59-Nsp15-HA have increased sgRNA and cellular RNA packaged into virions. We next assessed the degree of the selective packaging defect observed in rA59-PS- and rA59-Nsp15-HA virions (Fig. 2). To accomplish this goal, the levels of positive- and negative-strand sgRNA7 (+sgRNA7 and −sgRNA7, respectively) and positive-strand gRNA (gRNA) were determined using concentrated virus samples prepared by velocity centrifugation of infected-cell supernatants. In agreement with previous publications (8, 11), both rA59-PS- and rA59-Nsp15-HA virions had higher levels of packaged +sgRNA7 than gRNA compared to rA59WT virions (Fig. 2A). Furthermore, the ratio of +sgRNA7 to gRNA in rA59-PS- and rA59-Nsp15-HA virions was nearly equivalent to the intracellular ratio in rA59-PS- infected and rA59-Nsp15-HA-infected cells. This enrichment of +sgRNA7 to gRNA in rA59-PS- and rA59-Nsp15-HA virions contrasts with rA59WT, where the +sgRNA7 to gRNA ratio was dramatically decreased compared to the intracellular ratio (Fig. 2A). An increased ratio of −sgRNA7 to gRNA was also observed in rA59-PS- and rA59-Nsp15-HA compared to rA59WT virions (Fig. 2B) but was not equivalent to the intracellular ratio of these RNAs. Finally, these increases in sgRNA7 incorporation had little effect on the levels of gRNA (Fig. 2C), as neither rA59-PS- nor rA59-Nsp15-HA virions had more than an ~2-fold decrease in packaged gRNA compared to rA59WT virions.
In retroviruses, the loss of gRNA packaging results in robust packaging of cellular RNA (10). To determine the extent of cellular RNA incorporation by MHV selective packaging mutants, we measured the ratio of hypoxanthine-guanine phosphoribosyltransferase (HPRT) RNA to gRNA in concentrated virus and intracellular RNA samples. We found a significant increase in the HPRT-to-gRNA ratio in both rA59 PS/H11002 and rA59Nsp15-HA (Fig. 2D) compared to rA59WT virions. However, the ratio of HPRT to gRNA in rA59PS/H11002 and rA59Nsp15-HA virions was not equivalent to the intracellular ratio of HPRT to gRNA. These data establish that rA59PS/H11002 and rA59Nsp15-HA were similarly defective in selective packaging and suggest that cellular RNA can be packaged into virions at increased levels in selective packaging mutants.

rA59PS induces increased levels of IFN-I in BMMs. Because the two selective packaging mutants have similar phenotypes in vivo and similar packaging defects, we utilized only rA59PS and rJHMVPS to further investigate the mechanism of PS-mediated virulence. The elevated levels of sgRNA, especially −sgRNA, in virions may act as pathogen-associated molecular patterns (PAMPs), which may be sensed during entry, replication, or packaging of PS mutants. To initially test this hypothesis, we utilized bone marrow-derived macrophages (BMMs) because they are easily culturable, are highly susceptible to MHV, and are known to express IFN-I following infection (15).
BMMs infected with rA59PS/H11002 had an ~2- to 3-fold increase in beta interferon (IFN-β) at 12 h postinfection (hpi) compared to rA59WT-infected BMMs (Fig. 3A). The increased IFN-I response was not due to increased replication of rA59PS/H11002 as rA59PS/H11002 infection resulted in a small (~1.5- to 2-fold) reduction in gRNA (Fig. 3B). In agreement with previous reports, we found that IFN-I production was dependent on MDA5 for both rA59WT and rA59PS/H11002 (Fig. 3C). Furthermore, there was still a small difference (~1.5-fold) in replication of rA59PS/H11002 in MDA5/H11002 cells (Fig. 3D). These results indicate that alteration of the packaging signal in MHV can lead to an enhanced innate immune response and perhaps a modest defect in replication in primary cells.

**IFN-I signaling is required for rJHMVPS- attenuation.** Increased levels of IFN-I in BMMs following rA59PS- infection and the well-established protective role of IFN-I signaling in MHV infection (16) suggested that IFN-I signaling would be important for rA59PS- or rJHMVPS- attenuation. Since rJHMV was more virulent than rA59, rJHMVPS- was used to test this hypothesis. IFNAR/H11002 mice were infected with rJHMVPS- or rJHMVWT and monitored for morbidity and mortality for 14 dpi. rJHMVWT-infected IFNAR/H11002 mice had substantial weight loss and succumbed to infection earlier than rJHMVWT-infected wild-type mice (Fig. 4A and B compared to Fig. 1D and G). IFNAR/H11002 mice infected with rJHMVPS- at the standard dose of 3 × 10⁴ PFU succumbed to infection in a manner similar to infection with rJHMVWT (Fig. 4A and B). However, at a low dose of virus, 1 × 10³ PFU, the virulence of rJHMVPS- was not completely restored (Fig. S3). The increased levels of IFN-β in rA59PS- infected BMMs and rescued virulence of rJHMVPS- in IFNAR/H11002 mice suggest that IFN-I signaling plays an important role in the attenuation of rJHMVPS-. Additionally, other factors, such as delayed neuroinvasion, may also be important for the attenuation of PS- viruses.

Previous work has demonstrated protective roles for MDA5 and Toll-like receptor 7 (TLR7) during coronavirus infection (15, 17). Next, we utilized MAVS/H11002 mice to determine the role of MDA5-mediated signaling in attenuating rJHMVPS-. rJHMVWT caused a lethal infection in MAVS/H11002 mice, but rJHMVPS- remained attenuated, causing moderate weight loss and no decrease in survival (Fig. 5A and B). MHV induces significant levels of IFN-I in plasmacytoid dendritic cells via TLR7 signaling, contributing to protection (17). To determine the role of TLR7 signaling in protection, we infected...
TLR7\(^{-/-}\) mice with rJHMVs\(_{\text{PS}}\). Since the TLR7\(^{-/-}\) mice that were used were on a BALB/c background, we first confirmed that BALB/c mice were protected from rJHMVs\(_{\text{PS}}\)-mediated lethality. rJHMVWT infection caused complete lethality and substantial weight loss, while rJHMVs\(_{\text{PS}}\)-infected BALB/c mice had a phenotype similar to that observed in rJHMVs\(_{\text{PS}}\)-infected B6 mice, with increased survival and less weight loss (Fig. 5C and D). TLR7\(^{-/-}\) mice infected with rJHMVWT succumbed to infection, while we observed no increase in weight loss or lethality after infection with rJHMVs\(_{\text{PS}}\) (Fig. 5E and F). These results suggest that neither signaling through MAVS nor that through TLR7 is solely responsible for attenuation of packaging signal mutants and suggest that initiation of IFN-I signaling occurs via alternate sensors or is multifactorial.

While neither TLR7 nor MAVS signaling was solely responsible for rJHMVs\(_{\text{PS}}\) attenuation, it is possible that a protective level of IFN-I was still produced in infected MAVS\(^{-/-}\) or TLR7\(^{-/-}\) mice. IFNAR\(^{-/-}\) mice are incapable of IFN-I signaling and therefore lack expression of interferon-stimulated genes (ISGs). In contrast, MDA5\(^{-/-}\) mice infected with MHV had decreased but detectable levels of IFN-I, and the expression of many ISGs was not altered during MHV infection of MDA5\(^{-/-}\) mice (18). Next, to determine if IFN-I signaling during infection was responsible for protecting mice or if basal levels of ISGs were playing a significant role, we treated MAVS\(^{-/-}\) mice with an IFNAR-blocking antibody (19). In these experiments, IFNAR-blocking antibody was injected intraperitoneally 6 h prior to infection and again 3 days later. Control MAVS\(^{-/-}\) mice administered rat IgG remained protected from rJHMVs\(_{\text{PS}}\); conversely, MDA5\(^{-/-}\) mice administered IFNAR-blocking antibody displayed increased weight loss and succumbed to infection (Fig. 5G and H). These data suggest that IFN-I signaling during infection is important for the attenuation of rJHMVs\(_{\text{PS}}\).

**DISCUSSION**

Due to the largely unaltered growth of packaging mutants in vitro, the lineage A betacoronaviruses, specifically MHV, provide a unique opportunity to study the effect of viral packaging on virulence. This is not possible in both alphaviruses and retroviruses since packaging mutants have significant decreases in gRNA incorporation into virions and in infectivity (9, 10). Our data identify the lineage A betacoronavirus PS as a novel cis-acting virulence factor in MHV.

While the function of the lineage A betacoronavirus packaging signal is well understood in vitro, its relevance in vivo has not been addressed. The location of the stem-loop structure within the nsp15 locus allows these coronaviruses to selectively package gRNA and not coterminal sgRNA or host RNA. In this work, we quantified the packaging defect in rA59\(_{\text{Nop15-HA}}\) and rA59\(_{\text{PS}}\). The defects present in these two different PS mutants were strikingly similar, despite rA59\(_{\text{Nop15-HA}}\) maintaining the overall structure, the pentaloop, and one of the 2-nucleotide (nt) bulges, elements
hypothesized to be important for packaging (Fig. 1A and B). In agreement with previous publications (8), we found a significant increase in sgRNA packaging in both rA59PS and rA59Nsp15-HA virions. We further divided the sgRNA into +sgRNA and −sgRNA and found increased levels of both in the virions of mutant viruses compared to parental viruses. Furthermore, we found increased levels of host cellular RNA in both packaging mutant virions. Interestingly, while the ratios of +sgRNA7 to gRNA in the virions of both rA59PS and rA59Nsp15-HA were equivalent to the intracellular ratio, this was not true for ratios of −sgRNA7 or glyceraldehyde-3-phosphate dehydrogenase (GAPDH) to gRNA.

FIG 5 IFN-I signaling during rJHMVPS is not solely dependent upon either TLR7- or MAVS-mediated signaling. (A and B) Five- to 8-week-old male MAVS−/− mice were infected with 3 × 104 PFU of either rJHMVps− or rJHMVps+ by intranasal injection. Infected mice were monitored for survival (A) and weight loss (B) (rJHMVps−, n = 7; rJHMVps+, n = 12). (C and D) Five- to 8-week-old male BALB/c mice were infected with 1 × 105 PFU of either rJHMVps− or rJHMVps+ by intranasal inoculation. Infected mice were monitored for survival (C) and weight loss (D) (rJHMVps−, n = 5; rJHMVps+, n = 5). (E and F) Five- to 8-week-old male TLR7−/− (BALB/c) mice were infected with 1 × 106 PFU of either rJHMVps− or rJHMVps+ by intranasal inoculation. Infected mice were monitored for survival (E) and weight loss (F) (rJHMVps−, n = 5; rJHMVps+, n = 5). (G and H) Five- to 8-week-old male MAVS−/− (B6) mice were infected with 3 × 108 PFU of rJHMVps− by intranasal injection. At the time of inoculation and at 3 dpi, 1.5 μg and 0.5 μg of IFNAR-blocking antibody were administered, respectively, by the i.p. route to infected mice, which were monitored for survival (G) and weight loss (H) (rJHMVps−, n = 5; rJHMVps−, n = 7).
Packaging occurs through an interaction between the PS stem-loop and coronavirus structural proteins. A direct interaction between the M protein and the PS was previously reported and correlated with the packaging of nonviral RNA (20). It can be hypothesized that in the absence of M-PS interactions, packaging of RNA is dependent on N protein. Since N protein binds to RNA indiscriminately (5), levels of packaged RNA would be predicted to closely mimic those present in the intracellular environment. However, the ratio of both –sgRNA7 and GAPDH to gRNA was increased in virions but did not reach intracellular ratios, suggesting that RNA binding by N is not the sole determinant of RNA packaging in rA59PS and rA59_Nsp15-HA. RNA localization, secondary structure in other regions of the genome, or other proteins may influence RNA packaging. The relative lack of –sgRNA7 packaging may reflect its localization to membranous structures associated with replication while +sgRNA7 would be exported to the cytosol.

All packaging mutants tested were attenuated in mice. rJHMVPS and rJHMV_Nsp15-HA and rA59PS and rA59_Nsp15-HA were attenuated in the context of lethal encephalitis (Fig. 1) and hepatitis (see Fig. S1 in the supplemental material), respectively. It can be hypothesized that packaged sgRNA and/or host RNAs behave as PAMPs, eliciting an increased IFN-I response in rJHMVPS-infected and rJHMV_Nsp15-HA-infected cells. Consistent with this hypothesis, IFN-I signaling was essential for this attenuation in vivo (Fig. 3). Furthermore, BMMs infected with rJHMVPS expressed higher levels of IFN-I than rJHMVWT (Fig. 3A). These data indicate that packaging mutants are capable of inducing an increased IFN-I response, which may be protective in vivo. We investigated the two best-described sensors for IFN-I signaling during coronavirus infection, MDA5 and TLR7. We found that MAVS−/− and TLR7−/− mice were both resistant to rJHMVPS. These data indicate that neither TLR7- nor MAVS-mediated signaling was solely responsible for the attenuation of rJHMVPS. Furthermore, MAVS−/− mice treated with IFNAR-blocking antibody were more susceptible to rJHMVPS than control antibody-treated mice, suggesting that IFN-I was still responsible for rJHMVPS attenuation in these mice. Together, these data support a role for IFN-I signaling in rJHMVPS attenuation and suggest that TLR7 and MDA5 have a redundant function(s) or that other sensors/inflammatory pathways play a role in MHV packaging mutant attenuation.

The sgRNAs could act as PAMPs during the replication and/or entry of MHV packaging mutants. The presence of –sgRNAs in virions, albeit in small amounts, might be most important for the induction of IFN-I since negative-strand viral RNA is presumably not capped but, rather, contains a triphosphate at the 5′ end. This would facilitate recognition by the intracellular helical receptor RIG-I. Negative-sense sgRNA is expected to be present as double-stranded RNA (dsRNA), which could be detected by RIG-I, TLR3, and protein kinase R (PKR), in addition to MDA5. It should be noted that while our data suggest that IFN-I signaling and perhaps other changes in the innate immune response are largely responsible for rJHMVPS attenuation, defects in virus replication in cells in the infected brain may also contribute to attenuation. We observed no differences between the replication kinetics of rJHMVPS or rJHMV_Nsp15-HA and those of rJHMVWT in vitro (Fig. S1), but this does not preclude subtle differences in primary cells, such as BMMs (Fig. 3D).

In conclusion, our results demonstrate that the selective packaging of gRNA plays an important role in the virulence of MHV and probably other lineage A betacoronaviruses. This work uncovers a unique method by which an RNA virus shields its RNA products from the host to prevent an innate immune response. Selective packaging is conserved among coronaviruses and therefore is likely important for the virulence of most, if not all, coronaviruses. However, the localization of the packaging signal for most coronaviruses is unknown. Thus, an important goal for future research will be to identify the packaging signals in pathogenic human and animal coronaviruses and determine their roles in virulence.
**MATERIALS AND METHODS**

**Cell culture.** 17Cl-I cells, HeLa cells expressing MHV receptor carcinoembryonic antigen-related cell adhesion molecule 1 (CEACAM1a) (HeLa-MHVR), and baby hamster kidney cells expressing CEACAM1a (BHK-MHVR) were grown in Dulbecco’s modified Eagle’s medium (DMEM), with 10% fetal calf serum, 1% penicillin-streptomycin (Pen-Strep), 2% sodium bicarbonate, 2% L-glutamine, 1% nonessential amino acids, and 1% sodium pyruvate (D10). Low-serum medium is the same as D10 with 2% fetal calf serum (D2).

**Mice.** Pathogen-free C57BL/6 (B6) and BALB/c mice were purchased from Charles River Laboratories. B6 interferon alpha/beta receptor-knockout (IFNAR−/−) mice were obtained from the Jackson Laboratory, and Toll-like receptor 7-knockout (TLR7−/−) mice were obtained from Shizuoka Akira (Osaka University) via Westley Reeves (University of Florida). Mice were bred in the animal care facility at the University of Iowa. Animal studies were approved by the University of Iowa Animal Care and Use Committee (IACUC).

**Generation of recombinant viruses.** pBAC-rJHMVps−, and pBAC-rJHMVps/H11002 were created using pBAC-rJHMLIA, by Red recombination, and using the primers listed in Table S1 in the supplemental material. Using gBlock fragments with the corresponding mutations and listed primers, a PCR product was created with the HA or PS-containing product was transformed into competent GS1783 cells harboring the rJHMVIA pBAC. Choromaphenicol-positive (CML+) Kan− colonies were screened for intermolecular recombination by PCR and restriction enzyme digestion. Colonies with the correct intermediates were then used for intramolecular recombination of the homologous arms. Correct colonies were amplified, and the ISCE-I enzyme was induced with 1% arabinose for 2 h. The Lambda Red recombination enzymes were induced by heat shock at 42°C for 30 min, and cells were then incubated at room temperature for 4 h. CML− Kan− colonies were screened for correct final products by PCR, restriction enzyme digestion, and sequencing. pBACIAps− or pBACIA/H11002 was cotransfected with plasmid expressing N protein into BHK-MHVR cells using Lipofectamine 2000 (Fisher Scientific) according to the manufacturer’s protocol. Transfected BHK cells were collected at the time of maximal cytopathic effect and lysed by freeze-thawing. rJHMVps− and rJHMVps/H11002 were then passed 3 times to obtain working stocks. Two independent clones of the PS− virus (Fig. 1C), rJHMVps−, were isolated, and the sequence of each clone was confirmed. No coding mutations were present in either clone; however, rJHMVps−/ps+ did contain a synonymous mutation in nsp15 3′ of the PS. It is unlikely that this mutation significantly alters the secondary structure in this region, and this isolate behaved identically to rJHMVps−/ps−. rJHMVps−/ps+ was sequenced after passing with no mutations found.

To introduce the previously published PS− mutation (8) into rA59, the published mutations were introduced into the F plasmid of the rA59 system (21). The F plasmid was linearized, and homologous arms, with the PS− mutations, were added by PCR using the primers listed in Table S1. The linearized plasmid was recombined using the homologous arms and the In-Fusion cloning kit (Clontech) according to the manufacturer’s protocol. The PS− mutations were screened for by PCR and sequencing. The F plasmid and other fragments were digested with BsmBI and ligated together using T4 DNA ligase. Ligated DNAs and N protein transcript were transcribed in vitro using the mMessage mMachine T7 transcription kit (Fisher Scientific) according to the manufacturer’s protocol. BHK-MHVR cells were then transfected by electroporation with transcribed RNAs. Transfected BHK-MHVR cells were plated in D10 and monitored for cytopathic effects. Rescued recombinant viruses were collected, their titers were determined as described above, and they were termed rA59ps−. P0. rA59ps− P0 stocks were used to generate rA59ps−/ps−. P1 stocks by infecting 17Cl-I cells with virus at a multiplicity of infection (MOI) of 0.1 PFU/cell and collecting cells at times of maximal cytopathic effects. Titers of P1 stocks of rA59ps− and rA59ps+ were determined, and the stocks were utilized for all experiments with rA59.

**Tissue culture cell infection.** Unless otherwise noted, all infections were completed as follows. Semi-confluent monolayers of cells were infected on either 12-well or 24-well plates. Virus was adsorbed in DMEM at 37°C for 30 min at the indicated MOI. After adsorption, remaining virus was removed, and the cells were returned to growth medium for the indicated times.

**Mouse infection.** For rJHMV infections, 5- to 8-week-old male mice were sedated using isoflurane (Fisher Scientific). cDNAs for all RNAs except sgRNAs (see below) were prepared using Moloney murine leukemia virus (MMLV) reverse transcriptase with random hexamers according to the manufacturer’s instructions. Real-time quantitative PCR (RT-qPCR) was performed using Applied Biosystems QuantStudio 3 and PowerUp Sybr green master mix (Applied Biosystems). All primers used for qPCR are listed in Table S2. Target gene levels were normalized to hypoxanthine-guanine phosphoribosyltransferase (HPRT) by the threshold cycle (Ct) equation: ΔCt = CΔt − CΔt of the gene of interest − CΔt of HPRT. All results are shown as a ratio to HPRT calculated as 2−ΔΔCt.

**Selective viral RNA packaging.** Infected 17Cl-I supernatants were collected at 20 hpi, when maximal cytopathic effects were observed but before cell lysis. Supernatants were clarified by centrifugation at 1,000 × g and filtered through a 0.45-μm filter. Clarified supernatants were overlaid on 30% sucrose in Na-Tris buffer (100 mM NaCl, 10 mM Tris-Cl, pH 8.0) and centrifuged at 27,000 rpm for 4 h at 2,000 rpm at 4°C.
4°C. Pelleted virions were resuspended in DMEM. Viral RNA and cellular RNA from infected monolayers were isolated using Trizol (Fisher Scientific) according to the manufacturer’s protocol. cDNA for viral RNAs was generated using Superscript IV (Fisher Scientific) and strand-specific primers (Table S2) according to the manufacturer’s protocol, while cDNA for HPRT was generated using random hexamers as described above. Quantification for viral RNAs was performed utilizing a standard curve, which was generated from reverse-transcribed PCR-derived gRNA and sgRNA products. Relative quantification was performed using ΔCt, with normalization to gRNA. Data are presented as a ratio of +/– sgRNA7 RNA to gRNA, gRNA units per milliliter, or HPRT RNA to gRNA.

**Viral titers.** Viral titers were determined by infecting HeLa-MHVR cells as described above. Following rA99 infection, a 1:1 ratio of 1.2% agarose and gRNA. Following rRIMV infection, cells were returned to D2 medium for 16 h at 37°C. After incubation, a 1:1 mixture of 1.2% agarose and 0.1% crystal violet in 2% methanol. Following rJHMV infection, 24 h at 37°C. Following incubation, cell monolayers were fixed with 4% formaldehyde in phosphate-buffered saline (PBS) and stained with 0.1% crystal violet in 2% methanol. Following rJHMV infection, 24 h at 37°C. After incubation, a 1:1 mixture of 1.2% agarose and D2 with 1% neutral red was applied for 4 h at 37°C before counting plaques.

**BMM cultures.** Male and female 86 mice were euthanized, and bone marrow was isolated from each femur and tibia. The bone marrow was treated with ammonium-chloride-potassium lysing buffer for 30 s and then washed three times with RPMI containing 10% fetal calf serum. Bone marrow cells were then stained through 70-μm strainers and counted. Bone marrow cells (5 × 10^6) were plated in BMM medium (RPMI, 10% fetal calf serum, 10% L929 cell supernatant, 1% Pen-Strep, 1% sodium pyruvate, 1% l-glutamine) and cultured for 4 days. Following the 4th day, medium was changed daily until the day of experiment (day 7). BMMs were infected as described above at an MOI of 1 PFU/cell.

**Statistics.** Student’s unpaired t test or the Mann-Whitney U test was used to analyze differences in mean values between groups. All results are expressed as mean ± range or ± standard error of the mean where indicated. P values of ≤0.05 were considered statistically significant. Survival data were analyzed using Mantel-Cox tests (*, P < 0.05).

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at https://doi.org/10.1128/mBio.00272-18.

**FIG S1,** TIF file, 0.4 MB.

**FIG S2,** TIF file, 0.3 MB.

**FIG S3,** TIF file, 0.3 MB.

**TABLE S1,** DOCX file, 0.2 MB.

**TABLE S2,** DOCX file, 0.1 MB.

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