Dissection of Amino-Terminal Functional Domains of Murine Coronavirus Nonstructural Protein 3

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ABSTRACT
Coronaviruses, the largest RNA viruses, have a complex program of RNA synthesis that entails genome replication and transcription of subgenomic mRNAs. RNA synthesis by the prototype coronavirus mouse hepatitis virus (MHV) is carried out by a replicase-transcriptase composed of 16 nonstructural protein (nsp) subunits. Among these, nsp3 is the largest and the first to be inserted into the endoplasmic reticulum. nsp3 comprises multiple structural domains, including two papain-like proteases (PLPs) and a highly conserved ADP-ribose-1"-phosphatase (ADRP) macrodomain. We have previously shown that the ubiquitin-like domain at the amino terminus of nsp3 is essential and participates in a critical interaction with the viral nucleocapsid protein early in infection. In the current study, we exploited atypical expression schemes to uncouple PLP1 from the processing of nsp1 and nsp2 in order to investigate the requirements of nsp3 domains for viral RNA synthesis. In the first strategy, a mutant was created in which replicase polyprotein translation initiated with nsp3, thereby establishing that complete elimination of nsp1 and nsp2 does not abolish MHV viability. In the second strategy, a picornavirus autoprocessing element was used to separate a truncated nsp1 from nsp3. This provided a platform for further dissection of amino-terminal domains of nsp3. From this, we found that catalytic mutation of PLP1 or complete deletion of PLP1 and the adjacent ADRP domain was tolerated by the virus. These results showed that neither the PLP1 domain nor the ADRP domain of nsp3 provides integral activities essential for coronavirus genomic or subgenomic RNA synthesis.

IMPORTANCE
The largest component of the coronavirus replicase-transcriptase complex, nsp3, contains multiple modules, many of which do not have clearly defined functions in genome replication or transcription. These domains may play direct roles in RNA synthesis, or they may have evolved for other purposes, such as to combat host innate immunity. We initiated a dissection of MHV nsp3 aimed at identifying those activities or structures in this huge molecule that are essential to replicase activity. We found that both PLP1 and ADRP could be entirely deleted, provided that the requirement for proteolytic processing by PLP1 was offset by an alternative mechanism. This demonstrated that neither PLP1 nor ADRP plays an essential role in coronavirus RNA synthesis.
components that must cooperate in the integrated reactions required to produce capped and polyadenylated mRNAs and progeny genomes. Such colocalization would be most critical in the earliest stages of infection. Assembly of the replicase-transcriptase complex also induces extensive remodeling of intracellular membranes, resulting in the formation of double-membrane vesicles (DMVs), convoluted membranes, and other structures (6, 7). This transformation appears to involve, or else to coopt, the cellular process of macroautophagy, and it is triggered by the membrane-bound cohort of the replicase subunits. Expression of just nsp3, nsp4, and nsp6 has been shown to be necessary and sufficient to generate DMVs and the other membrane structures seen during infection (8). However, it is not currently clear whether membrane rearrangements are brought about by the virus specifically for the purpose of RNA synthesis or if, instead, they are by-products of the developing struggle between the incoming virus and host cellular defenses (9).

Among the components of the replicase, the first to be inserted into the ER is nsp3, via membrane-spanning segments near its carboxy terminus (10, 11). nsp3 is also, by far, the largest of the replicase subunits, having multiple structural domains (12, 13), all of which are oriented on the cytoplasmic side of the ER. Some of these modules are present, and are often well conserved, across all genera of coronaviruses. At the amino terminus of nsp3 is a ubiquitin-like domain (Ub1) (14) that is connected to a highly acidic (Ac) region of variable size (Fig. 1). In previous work, we discovered that MHV infection requires a critical interaction between Ub1 and the viral nucleocapsid (N) protein (15). Genetic, biochemical, and biophysical evidence has mapped this interaction to the serine- and arginine-rich region that falls between the two RNA-binding domains of N protein (15, 16). Moreover, we have shown that there exists a complete correspondence between the Ub1-N interaction and the requirement, common to all coronaviruses, for N protein to stimulate the infectivity of transfected genomes for the virus.

Three other nsp3 domains are common to all coronaviruses. The PLPs harbored by nsp3 are responsible for the first three events of polyprotein processing (Fig. 1). In MHV nsp3, there are two of these modules: PLP1 performs the nsp1-nsp2 and nsp2-nsp3 cleavages, while PLP2 carries out the nsp3-nsp4 cleavage (18, 19). Many, but not all, PLPs also possess a deubiquitinase activity, which acts to counter host innate immunity (20). Additionally, it has been suggested that PLPs may have some role other than proteolysis in replicase function (21). Between PLP1 and PLP2 in MHV (or else upstream of the sole PLP of some coronaviruses) there is a macrodomain with ADP-ribose-1-phosphatase (ADRP) and poly(ADP-ribose)-binding activity. The ADRP activity of this module has been shown to be dispensable in tissue culture (22, 23), but a role in viral RNA synthesis was proposed for the macrodomain, based on its nucleic acid-binding properties (24, 25). Finally, at the carboxy terminus of nsp3, there is a well-conserved region that has been dubbed the Y domain (26). Although nothing is known about its purpose, a particularly intriguing set of eight universally conserved cysteines and histidines occupies the amino-terminal 65 residues of this domain.

In order to better understand the functions of nsp3 in the replicase-transcriptase complex and how it contributes to the unique mechanism of coronavirus RNA synthesis, we began a dissection of MHV nsp3. Our effort was aimed at identifying those activities or structures in the huge nsp3 molecule that, like Ub1, are essential to the replicase. We also wanted to test the possibility of uncovering unknown roles of the better-characterized modules, such as the PLPs and the ADRP domain. In the present study, we found that both PLP1 and ADRP could be completely deleted, provided that the requirement for the proteolytic processing activity of PLP1 was offset by an alternative mechanism. This indicated that neither PLP1 nor ADRP plays an essential role in RNA synthesis, per se. Moreover, in the course of bypassing the need for PLP1 activity, we showed that viable mutants of MHV could be generated in which nsp1 and nsp2 were simultaneously eliminated.

**MATERIALS AND METHODS**

**Virus and cells.** Wild-type (WT), mutant, and revertant MHV strain A59 stocks were propagated in mouse 17 clone 1 (17C11) cells; plaque titrations and plaque purifications were carried out in mouse L2 cells. Monolayer cultures of both cell lines were maintained in Dulbecco’s minimal...
residues 86 to 127 of nsp1, His7, and 58 residues of foot-and-mouth disease virus (FMDV) VP1-2A (from GenBank accession number X00871). An epitope-tagged version of pA-SSN3, designated pA-SSN3HA, was built using the MiSeq 500-cycle v2 kit. Viral genome sequences were quantitated on a Qubit fluorometer (Invitrogen). Tagged libraries of DNA fragments were prepared with a Nextera XT kit (Illumina), and 2- by 250-bp paired-end sequencing was performed on a MiSeq Sequencer (Illumina). Viral mutants, RNA isolated from infected cell monolayers with the leader through nsp3 Ubl1 to PLP1) was used to replace the corresponding fragment of pA-WT. For pA-N3S, clone A of the N3S mutant, the BsiWI-BstBI fragment of pA-WT (running from the leader through nsp3 Ubl1) was replaced with a synthetic fragment encoding the nsp3 Ac region, purified for further analysis. To isolate revertants of the N3S mutant, 12 independent plaques obtained at 37°C were each used to start cultures that were grown at 37°C and thereafter serially passaged at 39°C in L2 cells at a low multiplicity of infection. When an increase in growth rate was noted, following 10 to 13 passages, titers of viral supernatants were determined on L2 cells at 39°C, infection. When an increase in growth rate was noted, following 10 to 13 passages, titers of viral supernatants were determined on L2 cells at 39°C, and a single larger plaque was chosen from each independent sample and purified for further analysis.

Genetic constructions were all made via manipulation of pBbar-A3 (17), a modified version of MHV cDNA clone A (27) that is here referred to as pA-WT. For pA-N3S, clone A of the N3S mutant, the BstBI-BstBI fragment of pA-WT (running from the leader through nsp3 Ubl1) was replaced with a fragment that was synthesized by PCR from overlapping oligonucleotides. The replacement fragment removed half of nsp1 and all of nsp2 and created 30 point mutations in 5’ RNA secondary structures, as described in detail in Results. For pA-SSN3, clone A of the SSN3 mutant, the BamHI-BstBI fragment of pA-WT (running from nsp1 codon 86 through nsp3 Ubl1) was replaced with a synthetic fragment encoding residues 86 to 127 of nsp1, His7, and 58 residues of foot-and-mouth disease virus (FMDV) VP1-2A (from GenBank accession number X00871). Plasmid pA-SSN3x3 was constructed identically to pA-SSN3, except that the NPGP consensus autoprocessing sequence was changed to NLAP (28).

An epitope-tagged version of pA-SSN3, designated pA-SSN3HA, was built in two steps. First, a synthetic fragment encoding the nsp3 Ac region, followed by FLAG and hemagglutinin (HA) tags, was inserted between the Sall and SacI sites binding the deleted Ac region of the previously described pA3Ac2, which had been used to generate the ΔAc2 mutant (17). Then the BstBI-Spel fragment of this intermediate (running from nsp3 Ubl1 to PLP1) was used to replace the corresponding fragment of pA-SSN3. Four derivative plasmids were obtained from pA-SSN3HA. One, pA-SSN3HAΔP3, was generated by a two-step PCR that replaced the SacI-Spel fragment (running from the start to the middle of nsp3 PLP1) to create the PLP1 catalytic mutation C289A. For the initial PLP1 deletion construct, pA-SSN3HAΔP3, the SacI-BstX1 fragment encompassing PLP1 was removed from pA-SSN3HA, and following the creation of blunt ends, the vector was reclosed. Oligonucleotides for PCR and DNA sequencing were obtained from Integrated DNA Technologies. The overall compositions of constructed plasmids were confirmed by restriction analysis, and all ligation junctions and regions resulting from PCR amplification were verified by DNA sequencing.

Viral growth kinetics. To measure growth kinetics, confluent monolayers of 17Cl1 cells (75 cm²) were inoculated at a multiplicity of 1.0 PFU per cell for 2 h at 37°C, with rocking every 15 min. Following the removal of inocula, monolayers were washed three times with phosphate-buffered saline (PBS), and incubation was continued in fresh medium at 37°C. Sample aliquots of medium were withdrawn at various times from 2 to 48 h postinfection, and infectious titers were subsequently determined in L2 cells.

Northern blotting. RNA was extracted from infected cell monolayers with TRI-Reagent (Zymo) according to the manufacturer’s instructions. Northern blotting of intracellular RNA was performed as previously described in detail (29) by using a PCR-amplified probe corresponding to the 3’-most 539 nucleotides of the N gene and the entire 3’ untranslated region (UTR) of the MHV genome. The probe was labeled with an AlkPhos Direct kit, and blots were visualized using CDP-Star detection agent (GE Healthcare).

In vitro translation. Capped mRNAs were produced with T7 RNA polymerase (mMessage mMACHINE; Ambion) by run-off transcription of BatZ17I-truncated pA-WT or SpeI-truncated pA-N3S, pA-N3SA1, pA-N3SA2, pA-SSN3, or pA-SSN3x3. Synthetic transcripts (0.46 μg) were translated in 17 μl of rabbit reticulocyte lysate (Ambion) labeled with [35S]methionine (MP Biomedical); protein products were analyzed in 10% polyacrylamide gels by SDS-PAGE, followed by fluorography. Vector pA-N3SA1 was derived from pA-N3S by removal of the BsiWI-NcoI fragment and religation of the plasmid following the creation of blunt ends. The same strategy was used to make pA-N3SA2 by removal of the Apal-Ncol fragment.

Western blotting. Sets of lysates were prepared from 75-cm² confluent monolayers of L2 cells that were mock infected or infected with wild-type and mutant MHV at either a high or a low multiplicity of infection at 37°C. At 8 to 10 h postinfection (for infections begun at a multiplicity of 1 PFU per cell) or 17 to 27 h postinfection (for infections begun at a multiplicity of 0.01 PFU per cell), monolayers were washed twice with PBS and then lysed by the addition of 500 μl of 50 mM Tris-HCl (pH 8.0), 150 mM NaCl, 1.0% Nonidet P-40, 0.7 μg/ml aprotinin, and 0.5 mg/ml Pefabloc SC (Roche). Lysed cells were held on ice for 15 to 30 min and then clarified by centrifugation. For immunoprecipitations, lysates were cleared with preimmune antiserum and were then incubated for 1 h at 4°C with anti-nsp3 antibody D3 (30) or VU164 (31), generously provided by Susan Baker (Loyola University Chicago, Maywood, IL) or by Mark Denison (Vanderbilt University, Nashville, TN), respectively. Samples were incubated for an additional 1 h at 4°C with a 75% slurry of nProtein A Sepharose (GE Healthcare) in lysis buffer. Sepharose beads were collected by centrifugation, washed three times with lysis buffer, and used directly for SDS-PAGE sample preparation. Lysate samples or immunoprecipitates were separated by SDS-PAGE through 7.5% or 4-to-15% gradient polyacrylamide gels, with prestained protein markers (NEB) in flanking lanes, and were transferred to polyvinylidene difluoride membranes. Blots were probed with one of the following: anti-HA monoclonal antibody (MAb 12CA5; Roche), anti-nsp1 antibody VU221 (32), or anti-nsp8 antibody VU124 (33); the last two antibodies were also gifts from Mark Denison. Bound primary antibodies were visualized using a chemiluminescent detection system (ECL or West Dura; Pierce).
RESULTS

Construction of a mutant containing nsp3 as the first subunit of the replicase-transcriptase complex. To begin to dissect the amino-terminal domain structure of nsp3, it was first necessary to disentangle nsp3 from its obligation to process upstream nsp subunits. The PLP1 module of MHV nsp3 carries out the nsp1-nsp2 and nsp2-nsp3 cleavage events of replicase-transcriptase polyprotein maturation (Fig. 1). Our first strategy to obviate these cleavages was to create a mutant, designated nsp3-start (N3S), in which polyprotein translation initiated with nsp3, while most of the preceding coding region was deleted. The major constraint on the construction of the N3S mutant was the need to preserve genomic elements that participate in viral RNA synthesis. The 5’ ends of coronavirus genomes, including the 5’ UTR and part of the nsp1 coding region, contain multiple cis-acting RNA structures that are critical for replication and transcription (34). In MHV and other betacoronaviruses, the genus in which these structures have been most extensively studied, the genomic 5’ end contains a series of at least eight stem-loops (SL) (35–39), as well as a four-way junction formed by a long-range interaction between the regions upstream of SL5 (SL5) and downstream of SL7 (40) (Fig. 2A). Mutations or deletions in many of these elements have been found to debilitate or kill the virus or to abrogate defective interfering (DI) RNA replication.

We consequently designed the 5’ UTR of the N3S mutant to contain a modified version of the 5’-most 594 nucleotides of the MHV genome, thereby retaining some margin beyond the 467-nt minimal segment previously shown to be sufficient to support DI RNA replication (41, 42). The AUG codon that occurs at nucleotides 592 to 594 was fused to the beginning of the nsp3 coding region (nt 2707 of the wild-type sequence) to serve as the start codon for the replicase-transcriptase polyprotein (Fig. 2B); at the −4 position relative to the AUG codon, nt 588 was mutated to optimize the context for translation initiation (43). To ensure utilization of the newly created start codon, the original start codon for nsp1 and the other four in-frame upstream AUG codons were each knocked out by mutations at 2 or 3 positions. Additionally, six out-of-frame upstream AUG codons were each mutated at 1 or 2 positions. Further point mutations were made to restore any base pairs in RNA secondary structures that would be disrupted by the start codon knockouts. The RNA secondary structure shown in Fig. 2 is the lowest free-energy mfold structure (44) for wild-type MHV nucleotides 79 to 594 (with no imposed constraints to generate the long-range interaction). The mfold-predicted structure for the N3S mutant was verified to be identical to that of the wild type. The only upstream start codon that was not eliminated was that for the small upstream open reading frame (uORF) in SL4 (Fig. 2B). Similarly situated uORFs are found in the majority of coronavirus genomes, and in MHV there exists selective pressure for the uORF to be maintained (45).

Following multiple trials, a single isolate of the N3S mutant was obtained. Initially, the sequence of the 5’ genomic end of this recombinant was determined, confirming that it contained exactly the engineered 30 point mutations and 2,112-nt deletion (Fig. 2B) and, in particular, that the 5’-most AUG codon occurred at nt 592 to 594 (Fig. 3A). To verify the functionality of this relocated start codon, we translated synthetic mRNAs that were identical to the 5’ ends of the wild-type or N3S mutant genomes. These mRNAs were generated by runoff in vitro transcription of truncated pA-WT or pA-N3S, the first of the seven plasmids used for assembly of full-length wild-type or N3S mutant cDNAs, respectively (Fig. 3B). Additionally, we produced two N3S-related mRNAs from plasmids pA-N3SΔ1 and pA-N3SΔ2, which contained deletions in the 5’ UTR of pA-N3S. The mRNAs were translated in a reticulocyte lysate system, and [35S]methionine-labeled protein products were analyzed by SDS-PAGE. For pA-WT mRNA, a product corresponding to the entire nsp1 fused to a small fragment of nsp2 had a size consistent with its expected molecular mass of 40.2 kDa (Fig. 3C). Translation of pA-N3S mRNA yielded a partial nsp3 fragment with an apparent molecular mass of 46 kDa, somewhat larger than its calculated size of 35.9 kDa. However, since a product of identical mobility was obtained with pA-N3SΔ1 and pA-N3SΔ2 mRNAs, the first of which has a 5’ UTR of only 25 nt, this showed that translation of pA-N3S mRNA had started at the expected location. As anticipated for cap-dependent protein synthesis, the efficiency of translation of the N3S-related mRNAs increased as the size of the 5’ UTR decreased. The aberrantly slow migration of the partial nsp3 protein was likely due to the highly acidic (Ac) region near the amino terminus of nsp3 (Fig. 1); the same mobility was seen for an almost identical nsp3 fragment that was initiated by a completely different mechanism (see below). A faint, slower-migrating band was observed for translated pA-N3S mRNA (Fig. 3C, open circle), but this same band was present for the 5’-UTR-deleted N3S mRNAs and it therefore could not have resulted from upstream initiation at a noncanonical start codon.

Together, these results demonstrated that the complete elimination of nsp1 expression coupled with deletion of the entirety of nsp2 does not abolish the viability of MHV in tissue culture. This represents the largest truncation of the replicase-transcriptase that has yet been made, although it must be noted that successful individual deletion of nsp2 in both MHV and SARS-CoV has already been well established (31). Combined with our prior work (17), this means that Ubl1, the amino terminus of nsp3, is the first functional domain of the replicase-transcriptase with respect to viral RNA synthesis. Nevertheless, because the N3S mutant had been very difficult to isolate, this raised the possibility that it needed to acquire one or more adaptive mutations in order to become fully viable. The determination of the complete genomic sequence of this mutant revealed that, indeed, it harbored two mutations in the nsp3 coding region. The first of these, E851G, was at the amino-terminal end of PLP2, near the Ubl2 domain. The second mutation, D1791V, fell in the carboxy-terminal Y domain. There were only three additional mutations in the N3S genome outside nsp3. One of these, T220I in nsp15, was at the boundary of a nonessential surface loop on this subunit, and we thus think it is not functionally relevant. Likewise, two mutations in the spike (S) protein, Q99H in the ectodomain and C1280S in the transmembrane domain, could conceivably affect growth but are highly unlikely to signify an interaction between S and nsp3.

The phenotype of the N3S mutant also indicated that it was conditionally impaired. Plaques of N3S were moderately smaller than wild-type plaques at 33°C and 37°C. However, at 39°C, the mutant formed plaques that were tiny compared to those of the wild type (Fig. 4A). To explore the basis for this deficiency, 12 cultures begun from individual plaques of N3S were passaged serially 10 to 13 times at 39°C in L2 cells. Titers of the viral supernatants were then determined at 39°C, and one revertant plaque originating from each culture was purified for analysis. Each cho-
FIG 2. Strategy for construction of the nsp3-start (N3S) mutant. (A) Schematics of the genomic 5' ends of wild-type MHV and the N3S mutant. Numbered RNA structures, stem-loops SL1 through SL7 (35, 36, 39) and SL8 (37), have been characterized previously. LR indicates a long-range interaction between the regions upstream of SL5 and downstream of SL7 (40). We have designated three structures predicted for the region downstream of SL8 as SLA, SLB, and SLC. Asterisks in the N3S genome show the positions of AUG start codons that were knocked out; open circles in the wild-type genome denote cleavage sites for PLP1. Note that the schematics do not accurately depict the relative lengths or extent of overlap of the 5' RNA secondary structures and the nsp1 ORF. (B) Details of the changes engineered in the N3S mutant to disrupt 5 in-frame start codons and 6 out-of-frame start codons upstream of the nsp3 ORF. Only structures that contain start codons are shown. Residue numbering begins from the 5' end of the wild-type genome; the N3S mutant has a deletion removing wild-type nucleotides 595 through 2706. Nucleotides of in-frame start codons are boxed; nucleotides of out-of-frame start codons are circled. In SL4, the start and stop codons of the uORF, denoted by stars, were unaltered in the N3S mutant. In SL5, the wild-type nsp1 start codon is labeled. N3S mutations, indicated by arrows, were made to knock out start codons, to preserve RNA secondary structure, or to optimize the context of the start codon that was juxtaposed to the start of the nsp3 ORF.

Among the revertants, those originating from the N3S mutant formed plaques that were substantially larger than those of the N3S mutant (two examples are shown in Fig. 4A), but none reached the size of wild-type MHV plaques at 39°C. For all 12 independent revertants, we sequenced the 5' end of each mutant (two examples are shown in Fig. 4A), but none reached the size of wild-type MHV plaques at 39°C. All 12 independent revertants sequenced the 5' ends of the leader through the start of the nsp4 gene, as well as the entire N gene and the 3' UTR. Only one revertant (rev12) had a nucleotide change (U476A) in the 5' UTR, which occurred in a bulge base of SLA (SLA) (Fig. 2B). Otherwise, no alterations were found in the 5' or 3' UTRs of any revertant. This suggested that the changes engineered in the original N3S mutant did not adversely affect the functioning of cis-acting RNA elements at either end of the genome.
In contrast, all revertants exhibited from one to three mutations in the nsp3 coding region, and these were gathered in three main loci (Fig. 4B). Multiple reverting mutations mapped to the amino-terminal Ubl1 and Ac domains. Three of these—A38V, A60T, and D98Y—were positioned within or close to the α2 helix of Ubl1 previously shown to interact with N protein (16). Moreover, A38V was identical to one of the reverting mutations originally seen in the N protein chimeric mutant that led us to discover the Ubl1-N interaction (15). Conversely, though, none of the N3S revertants contained changes in the serine- and arginine-rich region of N protein. An nsp3 mutation found in one revertant, L480I, mapped to the border between the PLP1 and ADRP domain; however, it is not currently clear why the unique architecture of the N3S mutant genome necessitates its acquisition over, and nsp2-nsp3 cleavage events, making use of the 2A peptide autotranslating element from the picornavirus foot-and-mouth disease virus (FMDV). Through an undefined mechanism, the FMDV 2A sequence effectively dictates termination followed by reinitiation of protein synthesis at two adjacent codons in a single ORF (46). Exploiting this device, we constructed a mutant, designated stop-start-NSP3 (SSN3), in which synthesis of the replicase-transcriptase polyprotein began with an amino-terminal segment of nsp1 that was separated from nsp3 by a 2A element (Fig. 5A). The fragment of nsp1 (amino acids 1 to 127) was chosen to preserve the same cis-acting genomic RNA structures as had been

FIG 3 Confirmation of the N3S mutant. (A) Segment of sequence of an RT-PCR product from total RNA purified from N3S-infected cells. The newly created nsp3 start codon is boxed. (B) Schematics of mRNAs generated by runoff in vitro transcription of truncated T7 vectors. Plasmids pA-WT and pA-N3S were those used to produce the 5′-most fragment of the full-length genomic cDNA for wild-type virus and N3S mutant virus, respectively. The length of the 5′ UTR is given above each mRNA. Deleted derivatives of pA-N3S reduced the size of the 5′ UTR of the encoded mRNA from 591 nt to either 25 nt (pA-N3SΔ1) or 249 nt (pA-N3SΔ2). The three pA-N3S-related plasmids were truncated with SpeI; pA-WT was truncated with BstZ17I. In the 5′ UTRs of mRNAs, the uORF is denoted by an open rectangle; asterisks represent start codon knockouts. The predicted molecular masses of translation products are indicated above the mRNAs. (C) In vitro-transcribed mRNAs were translated in a reticulocyte lysate system, and [35S]methionine-labeled protein products were analyzed by SDS-PAGE, followed by fluorography. Mock, control without added mRNA; [nsp3], nsp3 partial product from pA-N3S-related runoff mRNAs; nsp1-[nsp2], nsp1-partial nsp2 product from pA-WT runoff mRNA. The open circle denotes an artifactual band obtained with pA-N3S-related runoff mRNAs. The [nsp3] and nsp1-[nsp2] proteins contain 4 and 10 methionine residues, respectively.
A related mutant, SSN3x (Fig. 5A), that was modified by two sequential derivative mutants were acquired, we were unable to obtain a His7 epitope tag (data not shown).

nsp2, irrespective of the presence of either the 2A element or the constructed mutants that contain large deletions in both nsp1 and nsp3. This was expected to be tolerated, because the first 18 residues of MHV Ubl1 are unstructured and do not contribute to the interaction with N protein (16).

We first implemented this design with the minimal 20-amino-acid FMDV 2A peptide, but no viable virus was obtained. Similarly, inclusion of a homolog of the 2A peptide, from _Thosea asigna_ picornavirus, appeared to be only partially effective, since recovered virus was very impaired. Amino-terminal extensions of the 2A element have been shown to greatly increase stop-start efficiency, consistent with the notion that the stretch of polypeptide occupying the ribosomal exit tunnel causes or contributes to this anomalous translational mechanism (47, 48). Accordingly, the final version of the SSN3 construct incorporated a 20-amino-acid FMDV 2A peptide plus 38 upstream residues from its native polypeptide sequence, preceded by a His7 epitope tag (Fig. 5A). Three independent isolates of the SSN3 mutant were obtained, all having the same phenotype, and one was chosen for further analysis. The sequence of this virus was determined from the 5′ end of the genome through the PLP1 domain and was confirmed to be exactly as expected, with no additional mutations. Plaques formed by the SSN3 mutant were only slightly smaller than wild-type plaques at 37°C and 39°C (Fig. 5). However, at 33°C, we consistently noted heterogeneity in plaque size, with the sporadic appearance of much smaller plaques. The basis for this growth characteristic at 33°C is unknown, but we have observed it for all representative revertants of N3S at 39°C.

Contrary to the direct fashion in which the SSN3 and subsequent derivative mutants were acquired, we were unable to obtain a related mutant, SSN3x (Fig. 5A), that was modified by two changes in the FMDV 2A sequence known to abolish autoprocessing (28). In four separate trials, each with multiple independent samples and robust positive controls, transfected SSN3x full-length RNA produced no signs of infection, indicating that the mutated 2A sequence was lethal. Since the only difference between the SSN3 and SSN3x constructs was the absence of a functional stop-start element in the latter, this strongly suggests that a permanent blockage of the amino terminus of nsp3 results in a non-functional replicase-transcriptase.

To seek further evidence for the relative efficacies of the SSN3 and SSN3x stop-start elements, we _in vitro_ translated synthetic mRNAs that were identical to the 5′ ends of these mutant genomes. Translation of pA-SSN3 mRNA gave rise to two discrete products, the smaller of which was exactly the size expected (21.5 kDa) for the nsp1 fragment fused to the FMDV 2A element (Fig. 5C). The larger pA-SSN3 product, corresponding to the processed nsp3 fragment, migrated more slowly than expected for a 39.5-kDa protein. However, its mobility was identical to that of the product obtained by translation of pA-N3S1 mRNA, which, because of its minimal 5′ UTR, had served to produce a size standard for this nsp3 fragment (see Fig. 3C). (The nsp3 fragments encoded by pA-SSN3 and pA-N3S1 differ only at their amino-terminal residues, proline and methionine, respectively.) In contrast to pA-SSN3, translation of pA-SSN3x mRNA yielded just a single, larger protein with an apparent molecular mass of 65 kDa, bigger than the expected 57.4 kDa by the same differential observed for the other nsp3 products. The total absence of this unprocessed product upon translation of pA-SSN3 mRNA indicated that there was complete efficiency of termination by the incorporated 2A element. Moreover, the relative intensities of the bands for the nsp3 and nsp1-2A fragments (allowing for their different methionine contents) showed that reinitiation was also highly or entirely efficient. Conversely, the lack of production of either the nsp3 or

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**FIG 4** Revertants of the N3S mutant. (A) Plaques of the N3S mutant compared with those of isogenic wild-type virus at 33°C, 37°C, or 39°C, and plaques of two representative revertants of N3S at 39°C. Plaque titrations were carried out on L2 cells; monolayers were stained with neutral red at 72 h postinfection and were photographed 18 h later. (B) Locations of nsp3 mutations in 12 independent revertants of the N3S mutant. The two nsp3 mutations in the original N3S mutant, E851G and D1791V, are shown above the schematic.
nsp1-2A fragment upon translation of pA-SSN3x mRNA demonstrated that the mutations in the SSN3x construct effectively eliminated autoprocessing by its 2A element. Taken together, these results allowed us to conclude that the SSN3 construct enabled normal initiation of nsp1 with equimolar production of mature nsp3, thereby providing a suitable platform for further dissection of nsp3. The PLP1 and ADRP domains of nsp3 are entirely dispensable. Having eliminated the need for the two processing events carried out by PLP1, we were thus able to explore whether this module is required for other roles in the replicase-transcriptase complex. Toward this end, the SSN3 mutant was rebuilt with FLAG and HA epitopes inserted in tandem immediately downstream of the Ac region, yielding a construct designated SSN3HA (Fig. 6A). A PLP1 catalytic knockout, SSN3HAmutP, was then created by mutating an active site residue that has been shown to be essential for MHV PLP1 activity in vitro (18) and in vivo (19). To minimize the probability of spontaneous reversion of this mutation, C289A, we changed all three bases of the wild-type codon (UGU to GCC). Multiple independent isolates of both the SSN3HA and SSN3HAmutP viruses were readily obtained. As no phenotypic differences were noted among them, one isolate of each was chosen for further analysis. Sequence determination of the original SSN3 mutant at 33°C, 37°C, and 39°C (examples at 37°C are shown in Fig. 6B). This showed that, as anticipated, the knockout of PLP1 catalytic activity to have little or no effect in the SSN3 background, based on the prior demonstration of the viability of a PLP knockout mutant and its partial rescue by simultaneous small deletions in the nsp1-nsp2 end of the genome through most of nsp3 confirmed the presence in each of exactly the engineered composition with no extraneous mutations. Plaques formed by the SSN3HA and SSN3HAmutP viruses were identical to each other and to those of the original SSN3 mutant at 33°C, 37°C, and 39°C (examples at 37°C are shown in Fig. 6B). This showed that, as anticipated, the epitope tags were fully tolerated in what is thought to be an unstructured linker between the Ac region and PLP1 (12). Likewise, we had expected the knockout of PLP1 catalytic activity to have little or no effect in the SSN3 background, based on the prior demonstration of the viability of a PLP knockout mutant and its partial rescue by simultaneous small deletions in the nsp1-nsp2 and nsp2-nsp3 cleavage sites (19).

We next designed deletions of PLP1 in the SSN3HA background (Fig. 6A). Because no structural information is yet avail-

FIG 5 Construction and characterization of the stop-start-nsp3 (SSN3) mutant. (A) Schematics of the genomic 5' ends of wild-type MHV and the SSN3 and SSN3x mutants. In the wild-type genome, open circles denote cleavage sites for PLP1. In the mutant genomes, the expanded segment details the insertion placed between the nsp1 fragment and the start of the nsp3 coding region, which contains a His7 epitope tag followed by a 58-amino-acid version of the FMDV 2A stop-start element (47). The autoprocessing site in the SSN3 sequence is denoted by an arrowhead. Two inactivating mutations in the SSN3x sequence are underlined; the inactivated autoprocessing site is denoted by x. Indicated below each genome are the predicted molecular masses of translation products produced by runoff in vitro transcription and translation of plasmids used to construct full-length genomic cDNAs. (B) Plaques of the SSN3 mutant compared with those of isogenic wild-type virus at 33°C, 37°C, or 39°C. Plaque titrations were carried out on L2 cells; monolayers were stained with neutral red at 72 h postinfection and were photographed 18 h later. (C) Plasmids pA-WT, pA-SSN3, and pASSN3x, used for the transcription. The mRNAs were translated in a reticulocyte lysate system, and [35S]methionine-labeled protein products were analyzed by SDS-PAGE, followed by fluorography. Mock, control without added mRNA; [nsp1]-2A and [nsp3], autoprocessed products from pA-SSN3 runoff mRNA; [nsp1]-2A-[nsp3], unprocessed product from pA-SSN3x runoff mRNA; nsp1-[nsp2], product from pA-WT runoff mRNA. RNA transcribed from SpeI-truncated pA-N3SΔ1/SpeI (see Fig. 3), translated as a control, encodes an nsp3 fragment identical to that of pA-SSN3/SpeI mRNA, except that it has a methionine instead of a proline at the amino terminus. The [nsp1]-2A, [nsp3], [nsp1]-2A-[nsp3], and nsp1-[nsp2] proteins contain 7, 3, 10, and 10 methionine residues, respectively.
able for a betacoronavirus PLP1, other measures were used to infer the limits of this domain. In the first mutant, SSN3\textsuperscript{HA}\textDelta P\textsubscript{1}, the extent of the deletion (amino acids 234 to 484) was based on the smallest expressed segments of MHV nsp3 that exhibited protease activity \textit{in vitro} (49, 50), as well as on the boundaries of the upstream Ac (15) and downstream ADRP (23, 51) domains. In the second mutant, SSN3\textsuperscript{HA}\textDelta P\textsubscript{2} (deletion of amino acids 274 to 460), the boundaries of the first deletion were moved inward, to the edges of the region of PLP1 that is most conserved among the lineage A betacoronaviruses. In addition, the more conservative downstream boundary preserved residue M475, a mutation of PLP1 that has very far-reaching deleterious effects in a particular temperature-sensitive mutant, BR\textsuperscript{H9004}P3, which has one-fifth of nsp3; the sum of the deletions in nsp3 has a molecular mass of 260 kDa was detected (Fig. 7A). nsp3 has a molecular mass of 260 kDa (or 264 kDa with the FLAG epitope tag insertion) and has been described as 210 kDa (30, 31).

FIG 6 Mutation and deletion of domains of nsp3. (A) Schematics of the genomic 5’ ends of the SSN3\textsuperscript{HA} recombinant and mutants derived therefrom. At the top is a partial diagram of the domains of nsp3, as described in the legend to Fig. 1. In each mutant, HA marks the position of the FLAG and HA epitope tags inserted at the end of the Ac region. The solid rectangle between the nsp1 fragment and the start of the nsp3 coding region represents the FMDV 2A translational stop-start element, as in Fig. 5. In SSN3\textsuperscript{HA}\textDelta P\textsubscript{3}, the point mutation in PLP1 is indicated. In SSN3\textsuperscript{HA}\textDelta P\textsubscript{1}, SSN3\textsuperscript{HA}\textDelta P\textsubscript{2}, and SSN3\textsuperscript{HA}\textDelta P\textsubscript{3}, numbers denote the first and last residues of nsp3 deletions. Residue numbers given are those of mature wild-type nsp3 and do not include the extra amino-terminal proline or the 17-amino-acid signal sequence. (B) Plaques of the SSN3, SSN3\textsuperscript{HA}, SSN3\textsuperscript{HA}\textDelta P\textsubscript{1}, SSN3\textsuperscript{HA}\textDelta P\textsubscript{2}, and SSN3\textsuperscript{HA}\textDelta P\textsubscript{3} mutants compared with those of isogenic wild-type virus at 37°C. Plaque titrations were carried out on L2 cells; monolayers were stained with neutral red at 72 h postinfection and were photographed 18 h later.

To corroborate the genetic compositions of the SSN3\textsuperscript{HA} virus and its derivatives, we examined the expression of different subunits of the replicase-transcriptase. Infected cell lysates were immunoprecipitated with an antibody to the amino-terminal 347 amino acids of nsp3 (antibody VU164) (31) and then probed by Western blotting with an antibody to the HA epitope tag that had been inserted at the end of the Ac domain (Fig. 6A). (The FLAG epitope tag planted at the same internal position, adjacent to the HA tag, did not work in Western blots.) For cells infected with the SSN3\textsuperscript{HA} and SSN3\textsuperscript{HA}\textDelta P\textsubscript{2} viruses, a protein with an apparent molecular mass of 260 kDa was detected (Fig. 7A). nsp3 has a calculated molecular mass of 222 kDa (or 224 kDa with the epitope tag insertion) and has been described as 210 kDa (30, 31).
However, we observed the same band for wild-type, SSN3, SSN3HA, and SSN3HAmutP viruses in direct Western blots with VU164 antisemur (data not shown). We therefore conclude that the apparent 260-kDa band is nsp3, not a precursor, and that its lower mobility is probably attributable to differences between the molecular mass standards used by us and by others. Consistent with this, a single band of 235 kDa was observed for the SSN3HAΔP3 mutant, again larger than the calculated molecular mass of 178 kDa (Fig. 7A). This finding confirms that mature nsp3 of SSN3HAΔP3 was stably produced and that it harbored a large deletion. All of the same protein species were observed when immunoprecipitations were performed with a different anti-nsp3 antiserum (antibody D3) (30), which had been raised against the amino-terminal 205 amino acids of nsp3 (Fig. 7B). Notably, nsp3 of the SSN3HAΔP3 virus was only weakly immunoprecipitated by D3 despite containing the complete region of nsp3 recognized by this antibody. This may indicate that the deleted nsp3 of this mutant is conformationally different from wild-type nsp3.

Western blotting with anti-nsp1 antibody revealed a 26-kDa protein for wild-type-infected cell lysates (Fig. 7C), consistent with the expected molecular mass of 27.4 kDa for nsp1. The same antiserum also detected the 21.5-kDa nsp1-2A fragment of the SSN3 mutant and all of its derivatives, even though this fragment contained only half of the nsp1 molecule. (The His epitope tag that had been engineered upstream of the FMDV 2A element [Fig. 5] did not work in Western blots.) This result demonstrated that there was efficient in vivo termination by the 2A element, as we had observed in vitro (Fig. 5C). Additionally, Western blots of infected cell lysates probed with anti-nsp8 antibody detected the same 21.6-kDa species in wild-type- and mutant-infected cell lysates. This showed that, at the peak of infection, nearly equivalent amounts of this downstream replicase subunit were produced, suggesting that reinitiation by the FMDV 2A element was also highly efficient in vivo.

To more completely assess the fitness of the SSN3HA and SSN3HAΔP3 mutants, we evaluated their growth and RNA synthesis capabilities relative to those of the wild type. In infections begun at a multiplicity of 1.0 PFU per cell (limited by the titers that could be achieved for stocks of SSN3HAΔP3), wild-type virus reached peak titers of $7.8 \times 10^7$ PFU/ml between 8 and 12 h postinfection (Fig. 8A). In contrast, the SSN3HA and SSN3HAΔP3 mutants reached maximal titers that were 1.6 and 2.0 log$_{10}$ lower than those of the wild type with somewhat delayed kinetics, peaking between 12 and 16 h postinfection. Data from a second, independent growth experiment were nearly identical to those shown in Fig. 8A. We also noticed that 17Cl1 cell monolayers infected with wild-type MHV exhibited extensive syncytia and cytopathic effect and were almost completely detached by 24 h postinfection. However, 17Cl1 monolayers infected with either of the two mutants underwent a period of partial detachment from 16 to 24 h postinfection but had recovered to roughly 80% confluence by 48 h postinfection. This recovery was not observed with L2 cell monolayers. The differential growth kinetics (Fig. 8A) make clear that the major detriment caused by the engineering of the SSN3-derived mutants was due to the loss of nsp1 and nsp2. The gap between the growth curves for wild-type virus and both the SSN3HA and SSN3HAΔP3 mutants was roughly equal to the sum of the reductions seen previously for an individual nsp1 carboxy-terminal deletion mutant (53) and an individual nsp2 deletion mutant (31). Comparatively, the removal of the PLP1 and ADRP domains in the SSN3HAΔP3 mutant had only a minor effect. Northern blotting of intracellular RNA harvested at the peak of infection revealed a corresponding quantitative drop in RNA synthesis by the SSN3HA and SSN3HAΔP3 mutants relative to that of the wild type (Fig. 8B). However, both mutants produced the same subgenomic RNA species in the same relative proportions as wild-type MHV. This showed that deletion of the PLP1 and ADRP domains of nsp3 did not qualitatively alter the ability of MHV to execute the characteristic pattern of coronavirus RNA replication and transcription.

**DISCUSSION**

Coronaviruses are unique among RNA viruses in the size and complexity of their RNA-synthetic apparatus. Additionally, coronaviruses are highly unusual among positive-strand RNA viruses in that their genomic RNA is only minimally infectious in tissue culture unless supplemented with a source of N protein. In previous work, we have shown that the latter requirement is due, at least...
partly and perhaps entirely, to a critical association between N protein and the amino-terminal Ubl1 domain of nsp3 (15, 17). Our working model is that the purpose of the Ubl1-N interaction is to establish a connection between the infecting viral genome and the replicase that is being translated from the genome. This would coordinate the dual roles of genomic RNA, which functions initially as an mRNA and then subsequently as a template for RNA synthesis. Since nsp3 is the first replicase subunit to become anchored in the membrane, its amino terminus would be an advantageous location to dock the distal end of the infecting nucleocapsid, which is not traversed by ribosomes. nsp3 is a huge multidomain protein (222 kDa in MHV) constituting more than a quarter of the mass of the replicase-transcriptase complex (Fig. 1). The structures of many of its constituent modules have been determined, principally for the SARS-CoV homologs (13, 14, 16, 51). However, it is less clear what roles are played by most of these domains, except for the well-characterized processing and deubiquitinating activities of the PLPs (18–20, 30, 49). In the current study, we initiated a dissection of MHV nsp3 aimed at distinguishing which parts of this molecule are directly involved in RNA synthesis, whether in an enzymatic or structural capacity, and which parts are dispensable. Our strategy proceeded from the amino terminus. For Ubl1, we have previously shown that deletion of residues 19 through 111 or mutation of charged residues in the surface loop between the B3 and B4 strands is lethal to the virus (17). These results underlined the importance of the Ubl1-N interaction. In contrast, we previously showed that the Ac region, which is hypervariable even among closely related coronaviruses, can be deleted with no apparent effect on viral phenotype (17).

To continue downstream from Ubl1 and Ac, it was necessary to uncouple nsp3 from the processing of nsp1 and nsp2. The most straightforward approach toward this end was to remove nsp1 and nsp2 and to initiate polyprotein translation with nsp3. We accomplished this in the N3S mutant through a combination of a large deletion and knockout of all remaining start codons, subject to the strict maintenance of potential cis-acting RNA elements embedded in the 5′-most 591 nucleotides of the MHV genome (Fig. 2). The relocation of the replicase start codon to a point downstream of the 5′ cis-acting RNA elements showed that there is no strict requirement that they overlap the replicase ORF, even though this overlap is common to the genomic organization of all alpha- and betacoronaviruses. It is noteworthy that the 5′ UTR created in the N3S mutant was similar in size to those of gamma- and deltacoronaviruses, which have 5′ UTRs ranging from 477 to 606 nt and do not encode a counterpart for nsp1 (54, 55). To our knowledge, the N3S mutant is the first complete knockout of nsp1 (in an alpha- or betacoronavirus) and the first example of a coronavirus in which nsp1 and nsp2 have been simultaneously eliminated. This result confirms and extends prior findings with MHV and SARS-CoV.

MHV nsp1 mutants were previously constructed with carboxy-terminal deletions of 118 (53) or 33 (56) amino acids. These viruses exhibited relatively minor growth defects in cell lines and in primary cells, although the latter virus was shown to be severely attenuated in the mouse host. Extensive recent study of nsp1 has revealed it to be a suppressor of host protein synthesis, both by stimulation of a ribosome-associated endonuclease and by direct inhibition of translation initiation (57, 58). It thus appears that the main role of nsp1 is to institute a favorable cellular environment.

FIG 8 Growth and RNA synthesis by replicase-transcriptase mutants. (A) Growth kinetics of wild-type, SSN3HA, and SSN3ΔP3 viruses. Confluent monolayers of 17C11 cells were infected at a multiplicity of 1.0 PFU per cell. At the indicated times postinfection, aliquots of medium were removed, and infectious titers were determined by plaque assay on L2 cells. (B) Northern blot of total RNA isolated from mock-infected 17C11 cells or cells infected at a multiplicity of 1.0 PFU per cell with wild-type virus, the SSN3HA mutant, or the SSN3ΔP3 mutant. RNA was isolated from infected cells at 8.5, 10, or 12 h postinfection, respectively, for wild-type, SSN3HA, or SSN3ΔP3 virus. MHV RNA was detected with a probe specific for the 3′ end of the genome. gRNA, genomic RNA; sgRNA, subgenomic RNA. The right panel is an overlay-exposure to allow visualization of the larger RNA species.
for the infecting virus. The viability of the N3S mutant rules out any obligatory role for nsp1 in coronavirus RNA synthesis. An earlier study found that certain amino-terminal point mutations in nsp1 were severely debilitating or lethal for MHV (53), but, in retrospect, this may be attributable to disruption of cis-acting RNA structures overlapping the first half of the nsp1 ORF. A very recent analysis of the replication of a DI RNA of bovine coronavirus, which is closely related to MHV, found an absolute requirement for an intact stretch of coding sequence near the amino terminus of nsp1 (35). This nsp1 peptide was proposed to provide a means for DI RNA to be recruited to the preformed replication complexes of the helper virus. Our results argue that, even though this scheme is exploited for DI RNA replication, it does not reflect a mechanism normally used in viral RNA synthesis.

Similarly to nsp1, nsp2 cannot play an essential role in the coronavirus replicate-transcriptase complex. For both MHV and SARS-CoV, it was previously shown that the coding region of nsp2 could be entirely excised, resulting in mutants with roughly 10-fold reductions in peak growth titer (31). Although the nsp2 deletion viruses had a quantitative (2-fold) drop in RNA synthesis relative to wild-type levels, they exhibited no qualitative alteration of RNA species. In contrast to many of the domains of nsp3, where homology can be found across the most divergent genera, nsp2 is very poorly conserved, even among coronaviruses in the same genus. It is therefore unlikely to be specifically involved in the mechanism of RNA synthesis but instead may act in an ancillary capacity during infection. Indirect evidence suggests a role for SARS-CoV nsp2 in the alteration of intracellular signaling (59).

The recovery of the N3S mutant required two adaptive mutations in nsp3, and additional mutations in nsp3 were necessary to allow growth of revertants at 39°C (Fig. 4). These conditions did not provide a strong basis for further dissection of the nsp3 molecule. Consequently, we devised a different strategy, in the SSN3 mutant and its derivatives, to get around the polyprotein processing duties of PLP1. The FMDV 2A element (with an extended upstream peptide) was used to separate a fragment of nsp1 from the amino terminus of nsp3 (Fig. 5A). This device was seen to operate efficiently in vitro and in vivo, both in termination after the partial nsp1 molecule and in reinitiation to produce nsp3 and downstream replicase subunits (Fig. 5C and 7A and B). A minimal (20-amino-acid) version of the 2A element from Thosea asigna picornavirus had been used previously to replace the nsp1-nsp2 cleavage site of the human alphacoronavirus 229E (HCoV-229E) (60). However, it was not clear how effectively it functioned. In the context of the SSN3 mutant, there appeared to be an absolute requirement for processing at the amino terminus of nsp3. The lethality of the nonfunctional 2A element in the SSN3x mutant (Fig. 5A) is a strong indication that, under some circumstances, uncleaved polypeptide upstream of nsp3 can obstruct access to the Ubl1 domain. Our observation is consistent with, although more drastic than, previous demonstrations of the deleterious effects caused by deletion of the nsp2-nsp3 cleavage site (19) or by fusion of an inefficiently cleaved reporter protein to the amino terminus of nsp3 (61).

Within the framework of the SSN3 mutant, we were then able to address the requirements for the PLP1 and ADRP domains of nsp3. The multiplicity and specificity of nsp3 PLPs vary. Some coronaviruses, like SARS-CoV and MERS-CoV, have a single PLP that efficiently carries out all of the first three cleavage events of replicase polyprotein processing. Others, like MHV, have two PLPs with strictly designated roles; MHV PLP2 cannot be reactivated to perform both PLP1 cleavages (62). Still other coronaviruses, like HCoV-229E, have two PLPs with flexibly overlapping specificities (60). For coronaviruses with two PLPs, PLP1 is thought to have evolved through duplication of PLP2 (26). However, while the PLP1s of alphacoronaviruses remain akin to PLP2s (or single PLPs), the PLP1s of lineage A betacoronaviruses form a separate class with respect to substrate binding site structure (63). In this regard, MHV PLP1 may be considered a feature particular to the lineage A betacoronaviruses, comparable to the nsp15-encoded packaging signal, the hemagglutinin-esterase, and accessory proteins 2a and 5a. As such, PLP1 might not be expected to perform an essential RNA synthesis function conserved across all coronaviruses.

We found that a constructed PLP1 catalytic mutant, SSN³H₄₅mutP, had a phenotype identical to that of its parent viruses, SSN3 and SSN³H₄₅ (Fig. 6). This result was anticipated, because Graham and Denison had previously demonstrated the viability of a similar MHV PLP1 catalytic mutant (19). That virus, although severely impaired, was partially rescued when combined with small deletions of both the nsp1-nsp2 and nsp2-nsp3 cleavage sites. Their finding suggested that an inactivated PLP1 could stericly hinder replicate function by nonproductively binding to one or both of its cleavage sites. The SSN3 mutant background could thus be seen as an extreme form of the previous cleavage site deletions, in this case one in which absolutely no vestige of either PLP1 binding site was preserved. The SSN³H₄₅mutP virus showed that PLP1 proteolytic activity was not required for a hypothetical essential function in addition to polyprotein processing. It also provided a positive control for further dissection of nsp3, i.e., an inability to delete PLP1 would have to have been attributed to something other than loss of proteolytic activity.

It remained feasible that the PLP1 domain could harbor enzymatic or structural functions critical for viral RNA synthesis, and these would not be affected by the catalytic knockout mutation. This prompted us to test the possibility of completely deleting PLP1. A previous study of the PLP1 of HCoV-229E proposed that an as-yet undefined nonproteolytic role of PLP1 might be mediated by interactions between its universally conserved zinc finger and other nsp’s or RNA (21). Such a prospect, however, is precluded by our isolation of the SSN³⁴Δ₃ mutant, in which both PLP1 and the adjacent ADRP domain were deleted (Fig. 6). The SSN³⁴Δ₃ mutant removed 418 amino acids, more than one-fifth of the nsp3 molecule, ruling out any critical secondary role for the PLP1 domain in MHV RNA synthesis. Notably, the deletion in SSN³⁴Δ₃ (spanning amino acids 234 to 651) encompassed M475, the residue that is mutated in Brts31, a conditionally lethal MHV mutant (52). Paradoxically, at the nonpermissive temperature, Brts31 is defective in processing carried out by the main protease, nsp5, but it is unhindered in processing by PLP1 and PLP2. Our results suggest that the Brts31 mutation is dominant negative, manifesting the acquisition of an aberrant function at the nonpermissive temperature rather than the loss of a normal function.

In contrast to PLP1, the ADRP macrodomain (previously called the X domain [26]) is highly conserved in all coronavirus genera. A similar domain also appears in replicase proteins of alphaviruses, hepatitis E virus, and rubella virus. Moreover, in one alphavirus, Sindbis virus, macrodomain mutations that impair viral RNA synthesis have been identified (64, 65). In (uninfected)
cells, ADRP activity comes into play downstream of the pathway of RNA splicing, and thus a virally encoded ADRP appeared a good candidate to operate in an undefined analogous reaction in coronavirus RNA synthesis (66). Nevertheless, the ADRP enzymatic activity of coronavirus macrodomains has a very low turnover (22, 51, 67). Also, if hydrolysis of ADP-ribose-1’-monophosphate is critical for the virus, it is not clear why host cytoplasmic macrodomain proteins are inadequate for this task. Such considerations raised the possibility that ADP-ribose-1’-monophosphate is not the only substrate for the ADRP (67) or is a surrogate for the relevant substrate, which is possibly mono-ADP-ribosylated protein (68, 69). More to the point, ADRP active-site mutants of HCoV-229E (22) and MHV (23, 68) were found to be unimpaired in growth in tissue culture, although the latter were avirulent with respect to hepatic pathogenesis or induction of encephalitis. The knockout of ADP-ribose-1’-phosphatase clearly shows that this activity is not essential for coronaviruses, but it leaves unaddressed other potential roles of the macrodomain that shows that this activity is not essential for coronaviruses, but it raised the possibility that ADP-ribose-1’-monophosphate is not the only substrate for the ADRP (67) or is a surrogate for the relevant substrate, which is possibly mono-ADP-ribosylated protein (68, 69). More to the point, ADRP active-site mutants of HCoV-229E (22) and MHV (23, 68) were found to be unimpaired in growth in tissue culture, although the latter were avirulent with respect to hepatic pathogenesis or induction of encephalitis. The knockout of ADP-ribose-1’-phosphatase clearly shows that this activity is not essential for coronaviruses, but it leaves unaddressed other potential roles of the macrodomain that is not affected by active-site mutations (24). Some structural and biochemical studies have suggested that poly(ADP-ribose) binding and poly(A)-RNA binding are more conserved macrodomain activities that are more likely to be important for viral replication (24, 25). However, the complete deletion of the ADRP domain in the SSN3ΔΔP3 mutant unequivocally rules out an essential role for poly(ADP-ribose) binding, poly(A) binding, or any as-yet-unknown activities of the macrodomain in coronavirus RNA synthesis.

In the current study, we have covered nearly half of the MHV nsp3 ectodomain (Fig. 1). In future work, we would like to extend the same methods employed here to dissect the remainder of the nsp3 molecule. In particular, PLP2 and its associated Ubl2 domain (70) have prospects of playing roles in RNA synthesis beyond pro teaseolytic processing. Also, the carboxy-terminal Y domain (22), with its universally conserved clusters of cysteine and histidine residues, presents an intriguing target for potential intermolecular interactions. In addition, it would be of great interest to determine whether infections by SSN3, SSN3HA, or other mutants created here lead to the formation of the full spectrum of DMVs and other membrane alterations observed during wild-type infection. Such studies may help to elucidate the relationship of these membrane rearrangements to viral RNA synthesis (9).

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