

# Genetic Evidence of a Long-Range RNA-RNA Interaction between the Genomic 5' Untranslated Region and the Nonstructural Protein 1 Coding Region in Murine and Bovine Coronaviruses

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Higher-order RNA structures in the 5' untranslated regions (UTRs) of the mouse hepatitis coronavirus (MHV) and bovine coronavirus (BCoV), separate species in the betacoronavirus genus, appear to be largely conserved despite an ~36% nucleotide sequence divergence. In a previous study, each of three 5'-end-proximal *cis*-acting stem-loop domains in the BCoV genome, I/II, III, and IV, yielded near-wild-type (wt) MHV phenotypes when used by reverse genetics to replace its counterpart in the MHV genome. Replacement with the BCoV 32-nucleotide (nt) inter-stem-loop fourth domain between stem-loops III and IV, however, required blind cell passaging for virus recovery. Here, we describe suppressor mutations within the transplanted BCoV 32-nt domain that along with appearance of potential base pairings identify an RNA-RNA interaction between this domain and a 32-nt region ~200 nt downstream within the nonstructural protein 1 (Nsp1)-coding region. Mfold and phylogenetic covariation patterns among similarly grouped betacoronaviruses support this interaction, as does cotransplantation of the BCoV 5' UTR and its downstream base-pairing domain. Interestingly, cotransplantation of the BCoV 5' UTR and BCoV Nsp1 coding region directly yielded an MHV wt-like phenotype, which demonstrates a cognate interaction between these two BCoV regions, which in the MHV genome act in a fully interspecies-compliant manner. Surprisingly, the 30-nt inter-stem-loop domain in the MHV genome can be deleted and viral progeny, although debilitated, are still produced. These results together identify a previously undescribed long-range RNA-RNA interaction between the 5' UTR and Nsp1 coding region in MHV-like and BCoV-like betacoronaviruses that is *cis* acting for viral fitness but is not absolutely required for viral replication in cell culture.

In positive-strand RNA viruses that replicate in the cytoplasm, genomic 5'-end-proximal RNA structures carry out several functions required for virus reproduction. In coronaviruses, these are thought to include (i) translation initiation, commonly presumed to occur by a canonical cap-dependent 5'-terminal ribosomal entry mechanism, to synthesize the replicase enzymes from open reading frame 1 (13, 14, 31); (ii) signaling an RNA-dependent RNA polymerase template switch during minus-strand synthesis at a heptameric transcription-regulatory sequence (UCUAAAC in the case of mouse hepatitis coronavirus [MHV] and bovine coronavirus [BCoV]) (Fig. 1A and B) for placement of a common leader on subgenomic mRNAs (sgmRNAs) (43, 48, 49, 55); (iii) encoding signals on the 3' end of minus-strand genomic RNA (the antigenome) for initiating synthesis of plus-strand genomic mRNAs and sgmRNAs (9, 43, 48, 55); (iv) possibly harboring signals that act in *trans* to initiate synthesis of nascent plus-strand genomes (46); (v) possibly directly influencing initiation of minus-strand synthesis at the 3' end of the genome (33); and (vi) harboring a genome packaging signal (10, 18). A mechanistic understanding of these events may aid in the development of therapeutic inhibitors of coronavirus replication.

In a previous study (22), we exploited the ~36% nucleotide (nt) sequence divergence between the 5' untranslated regions (UTRs) of BCoV and MHV, separate species in the betacoronavirus genus (15) (formerly classified as group 2a coronaviruses), to identify potentially common 5'-end-proximal *cis*-acting elements for replication of MHV. Despite the relatively high nucleotide sequence divergence, higher-order structures in the 5'-end-proximal genomic regions of these two viruses are largely conserved (Fig. 1A and B) (7, 10, 29, 33, 35, 36, 41, 42), although the nomen-

clature is not yet uniform, since stem-loops (SLs) II, III, and IV in Fig. 1 are respectively named stem-loops 3, 4 and 5 by the Leibowitz, Giedroc, and Olsthoorn laboratories (10, 29, 33, 35, 36). It was learned that the BCoV SLI/II domain (nt 1 to 84), which contains higher-order structures named SLI (now known to be comprised of stem-loops 1 and 2) (29, 33, 35, 36) and SLII, can functionally replace its MHV counterpart, as can the SLIII domain (nt 85 to 141) and the SLIV domain (nt 174 to 226). For function, SLIV required a 16-nt extension into the nonstructural protein 1 (Nsp1) coding region to form a "long" SLIV (Fig. 1A and B). In each of these substitutions, despite nucleotide differences of 37, 22, and 33%, respectively, the chimeric virus replicated for at least three passages without compensatory suppressor mutations appearing within the 5'-most 1,000 nt and 3'-most 500 nt of the genome. In contrast, a 30-nt inter-stem-loop domain in the MHV 5' UTR that resides between SLIII and SLIV (nt 141 to 170) could not be replaced by the 32-nt BCoV counterpart (nt 142 to 173) to directly yield viable virus after transfection of the chimeric RNA genome (22). However, the resulting chimera, B<sup>142-173</sup>/M, produced plaques and viable progeny after two blind cell passages in

Received 8 September 2011 Accepted 2 February 2012

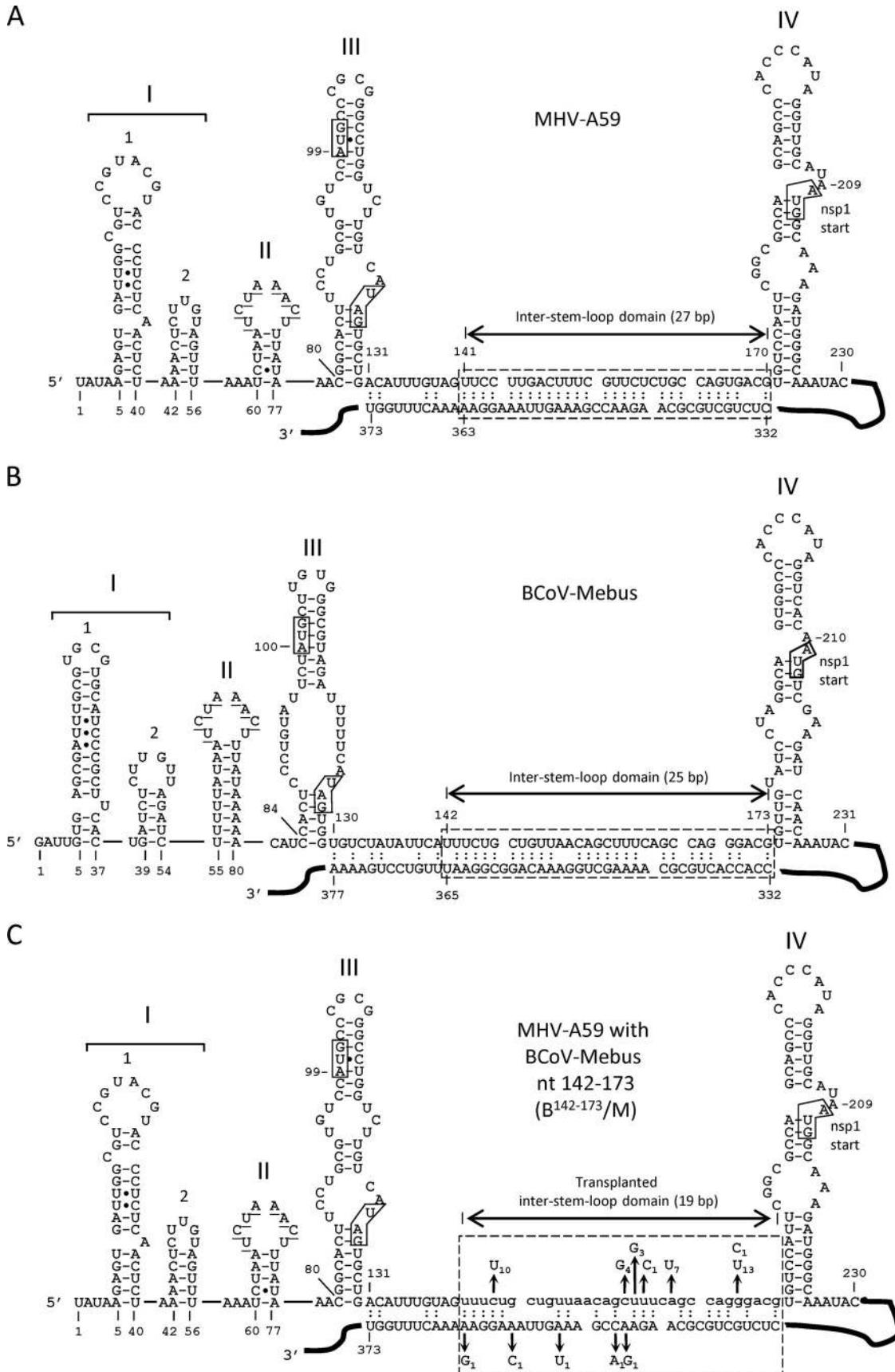
Published ahead of print 15 February 2012

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doi:10.1128/JVI.06265-11



one of four experimental attempts. These results suggested that there had been compensatory mutations within the surviving chimeras that enabled replication.

Here, we describe seven potential suppressor mutations that appeared within the transplanted 32-nt BCoV inter-stem-loop domain and five within a 32-nt partnering domain mapping ~200 nt downstream within the Nsp1 coding region and demonstrate the importance of these as cognate domains for replication in genetic reconstruction assays. The Nsp1 coding region is the first of 16 in the replicase gene (16, 21), and Nsp1 is an RNA binding protein with several postulated functions in coronavirus replication (5, 6, 23, 27, 28, 32, 39, 47, 56). Surprisingly, the 30-nt inter-stem-loop domain in the 5' UTR of the MHV genome can be deleted and viral progeny, although debilitated, are still produced. The results together identify a previously undescribed long-range *cis*-acting RNA-RNA interaction between the 5' UTR and Nsp1 coding region in the murine and bovine betacoronaviruses that appears to be important for viral fitness but is not absolutely required for virus replication in cell culture.

## MATERIALS AND METHODS

**Cells.** Delayed brain tumor (DBT) cells (24), mouse L2 cells (19), and baby hamster kidney cells expressing the MHV receptor (BHK-MHVR) (11, 12) were grown in Dulbecco's modified Eagle's medium (DMEM) supplemented with 10% defined fetal calf serum (FCS) (HyClone) and 20  $\mu$ g/ml gentamicin (Invitrogen). Cells were maintained at 37°C with 5% CO<sub>2</sub> for all experiments. BHK-MHVR cells were maintained in selection medium containing 0.8 mg/ml Geneticin (G418 sulfate; Invitrogen) (52).

**Viruses and GenBank accession numbers.** GenBank accession numbers for virus sequences analyzed are as follows: NC\_001846 for MHV-A59; NC\_006852 for MHV-JHM; NC\_012936 for Parker rat CoV; NC\_006577 for human coronavirus (HCoV)-HKU1; NC\_010327 for equine coronavirus (ECoV); NC\_005147 for HCoV-OC43; NC\_007732 for porcine hemagglutinating encephalomyelitis virus (PHEV); FJ415324 for human enteric coronavirus-4408 (HECoV-4408); U00735 for BCoV-Mebus; and NC\_004718 for SARS coronavirus, Toronto strain (SCoV-Tor2).

**RNA structure prediction and sequence alignments.** The Mfold program of Zuker (<http://www.bioinfo.rpi.edu/~zukerm/>) (38, 54) was used for RNA structure predictions. The long-range RNA-RNA base pairing patterns described below were revealed by folding nt 1 to 400 or nt 1 to 500. Sequence alignments were made with Vector NTI suite 8 software (Invitrogen).

**Reverse genetics system used for making BCoV/MHV chimeras and methods for analyzing viral progeny.** The reverse genetics system for MHV-A59, infectious clone MHV-A59-1000 (icMHV) (developed by R. Baric and colleagues [17, 52] and kindly provided by R. Baric), was used as previously described in detail for making 5'-end-proximal BCoV/MHV chimeras in the MHV-A59 background (22). Recombinant viruses described here were made by modifying fragment A of MHV-A59 as re-

ported previously (22) but with the appropriate primers for the mutations described below. All procedures for plasmid construction with icMHV DNA, expression of recombinant viral RNA, transfection of cells with infectious recombinant RNA by electroporation, and characterization of mutant progeny by virus titration and growth kinetics were carried out as previously described (22). Plaque morphology was determined on L2 cells after 60 h of growth and after crystal violet staining as described previously (22). Plaque sizes were identified as large (wild type [wt]) if they were  $\geq 2.5$  mm, medium if 1.5 to 2.5 mm, or small if  $\leq 1.5$  mm in diameter. Representative plaque images were taken with a Nikon digital camera and prepared using Adobe Photoshop CS software.

**Genome sequence analysis of chimera progeny.** Routinely, supernatant fluids from cells that first showed cytopathic effect (CPE) (either cells that had been transfected or cells that had been blind-passaged) were collected, and virus contained therein was referred to as virus passage zero (VP0) virus. When 80 to 100% of new DBT cells infected with VP0 virus showed CPE, intracellular RNA was extracted with TRIzol (Invitrogen), and the viral genome was sequenced by RT-PCR for the 5'-end-proximal nt 22 to 1093 and the 3'-terminal 500 nt (excluding the poly(A) tail). VP0 virus was then used to determine plaque morphology, and plaque-purified virus was used as starting material for growth kinetics on DBT cells and sequence analyses.

For analysis of the 5' nt 22 to 1093 of progeny virus genomes, extracted cellular RNA was reverse transcribed with Superscript II reverse transcriptase (Invitrogen) using primer MHV-1094(+) (5'CGATCAACGTGCCAAGCCACAAGG3'), which binds MHV genomic nt 1094 to 1117, and cDNA was PCR amplified with primers MHV-leader(-) (5'TATAAGAGTGATTGGCGTCCG3'), which binds nt 1 to 21 of the MHV antileader, or BCoV-leader(-) (5'GATTGTGAGCGATTGGCGTCCG3'), which binds nt 1 to 22 of the BCoV antileader, and MHV-1094(+). PCR products were gel purified (QIAEX II; Qiagen) prior to automated sequencing with primers MHV(261-284)(-) (5'CCATGGATGCTTCCGAACGCATCG3') and MHV(605-623)(+) (5'GTTACACAGGCAGACGCG3').

For analysis of the 3' 500 nt of progeny virus genomes, the same procedure was used except that total RNA was reverse transcribed using primer DI3(+) (5'CGGGATCCGTCGACACGCGTTTTTTTTTTTTTTT TTTTTT3'), which binds the poly(A) tail, and the cDNA was PCR amplified with primer MHV(30811-30830)(-) (5'GGATGGTGGTGCAGATGTGG3'), which binds the negative strand of virus nt 30811 to 30830, and primer DI3(+). Gel-purified PCR products were sequenced with primer MHV(30811-30830)(-).

**Northern analysis.** Northern analyses were done as previously described (25). Briefly, freshly confluent DBT cells in 25-cm<sup>2</sup> flasks ( $\sim 4 \times 10^6$  cells) were infected with wt or chimeric viruses at an MOI of 0.01 PFU/cell. At 20 h postinfection, intracellular RNA was TRIzol extracted, and 1/10 of the total RNA from one 25-cm<sup>2</sup> flask ( $\sim 60 \mu$ g RNA total per 25-cm<sup>2</sup> flask) was resolved by electrophoresis in a 1.0% agarose-formaldehyde gel at 150 V for 4 h. RNA was transferred to a Hybond N<sup>+</sup> nylon membrane (Amersham Biosciences) by vacuum blotting for 3 h followed by UV cross-linking. After prehybridization of the membrane with NorthernMax prehybridization/hybridization buffer (Ambion) at 55°C for 4 h, the blot was probed at 55°C overnight with 20 pmol ( $\sim 4 \times 10^5$  cpm/pmol) of the 5'-end  $\gamma$ -<sup>32</sup>P-labeled 3'-UTR-specific oligonucleotide

**FIG 1** 5' UTRs of MHV-A59 and BCoV-Mebus showing the intra-5'-UTR stem-loops, the inter-stem-loop domain, and a potential long-range RNA-RNA interaction between the 5' UTR and the Nsp1 coding region. (A) 5' UTR of MHV showing the Mfold-predicted long-range RNA-RNA interaction (hatched box). (B) 5' UTR of BCoV showing the Mfold-predicted long-range RNA-RNA interaction (hatched box). (C) 5' UTR of MHV in which the 32-nt BCoV inter-stem-loop domain (lowercase) replaces the 30-nt MHV counterpart. The hatched box identifies a potential long-range interaction and shows a summary of the potential suppressor mutations in 17 plaque-purified clones described in Table 1. The subscript for each mutation indicates its incidence among the 17 isolates. Note that without suppressor mutations, the number of base pairs within the window is 19, not 27 (as in MHV) or 25 (as in BCoV). Refolding enables the number of base pairs to increase as a function of suppressor mutations (see the text). In all panels, the underlined bases in SLII identify the transcription-regulatory sequence (UCUAAAC), and boxes in SLIII identify the start and stop codons for a short upstream open reading frame. The box in SLIV identifies the start codon for the Nsp1 coding region. Filled circles in SLI, SLII, and SLIII represent non-Watson-and-Crick base pairings, as determined by nuclear magnetic resonance studies (33, 36).

MHV(31094–31122)(+) (5' CAGCAAGACATCCATTCTGATAGAGA GTG3'), which binds MHV genomic nt 31094 to 31122. Probed blots were exposed to Kodak XAR-5 film at  $-80^{\circ}\text{C}$  for imaging, and images were prepared using Adobe Photoshop CS software.

## RESULTS

**Identification of seven mutations within the transplanted inter-stem-loop domain and five within the Nsp1 coding domain that potentially reconstitute base-pairing between the two domains in phenotypic revertants.** It was demonstrated previously that the 30-nt inter-stem-loop domain mapping between SLIII and SLIV in the MHV 5' UTR behaves as a species-specific *cis*-acting replication element, since the 32-nt homologue from the BCoV genome was essentially functionally incompatible with the MHV genome, as shown by the chimera B<sup>142–173</sup>/M (Fig. 1C) (22). Nucleotides 131 to 140, the 5'-terminal 10 nt of the entire sequence between SLIII and SLIV, were not included as part of the inter-stem-loop domain here or previously (22), because by some predictions it makes up part of SLIII in the MHV 5' UTR (29). Likewise, BCoV nt 130 to 141 were not included as part of the transplanted 32-nt inter-stem-loop domain. B<sup>142–173</sup>/M chimeric RNA containing the 32-nt transplanted BCoV domain during three transfection experiments was found to yield no viable progeny after four blind cell passages, but in a fourth trial, a mixture of small and wt-like large plaques were found after two blind cell passages (Fig. 2A, VP0 plaques). From this, six small plaques and 11 large plaques were selected, and each was passaged 10 times in DBT cells. Note that virus causing smaller plaques developed larger plaques and near-wt growth kinetics with passaging (Fig. 2). To determine what potential suppressor mutations might have caused the phenotypic changes within the transplanted 5' UTR, the adjacent Nsp1 coding domain, and the 3' UTR, viral genomic RNA isolated from cells infected at virus passages 1, 6, and 10 was sequenced through these regions. No 3'-end-proximal changes were found, and nucleotide changes within the 5'-end-proximal nt 22 through 1093 are summarized in Table 1.

The most striking result was a clustering of potential suppressor mutations within the transplanted 32-nt BCoV domain and within a 32-nt region mapping  $\sim 200$  nt downstream within the MHV Nsp1 coding region (summarized in Table 1 and schematically depicted in Fig. 1C and 3A). It was also striking that whereas there were fewer predicted base pairings initially between the two regions in B<sup>142–173</sup>/M RNA (19 bp) (Fig. 1C) than in Mfold-predicted wt MHV RNA (27 bp) (Fig. 1A), when predicted foldings included the putative suppressor mutations at virus passage 10 (Table 1), the number of base pairs within the window increased from 19 (Fig. 1C and 3A) to 25 (Table 1 and Fig. 3B). For example, when the combined suppressor mutations in S1 and S2 at VP10 were used, Mfold predicted the first potential folding pattern (potential folding 1) in Fig. 3B. Base pairings of 22 to 25 in revertants compare with 27 and 25 bp, respectively, for wt MHV and wt BCoV (Fig. 1A and B). These observations together led us to hypothesize that the suppressor mutations may reveal a specific long-range RNA-RNA interaction between the 5' UTR and the Nsp1 coding region that is important for virus replication.

**Mfold predicts base pairing between the inter-stem-loop domain and a region within the Nsp1 coding region of several MHV-like and BCoV-like betacoronaviruses.** After a potential base pairing between the two domains was first noted, as described

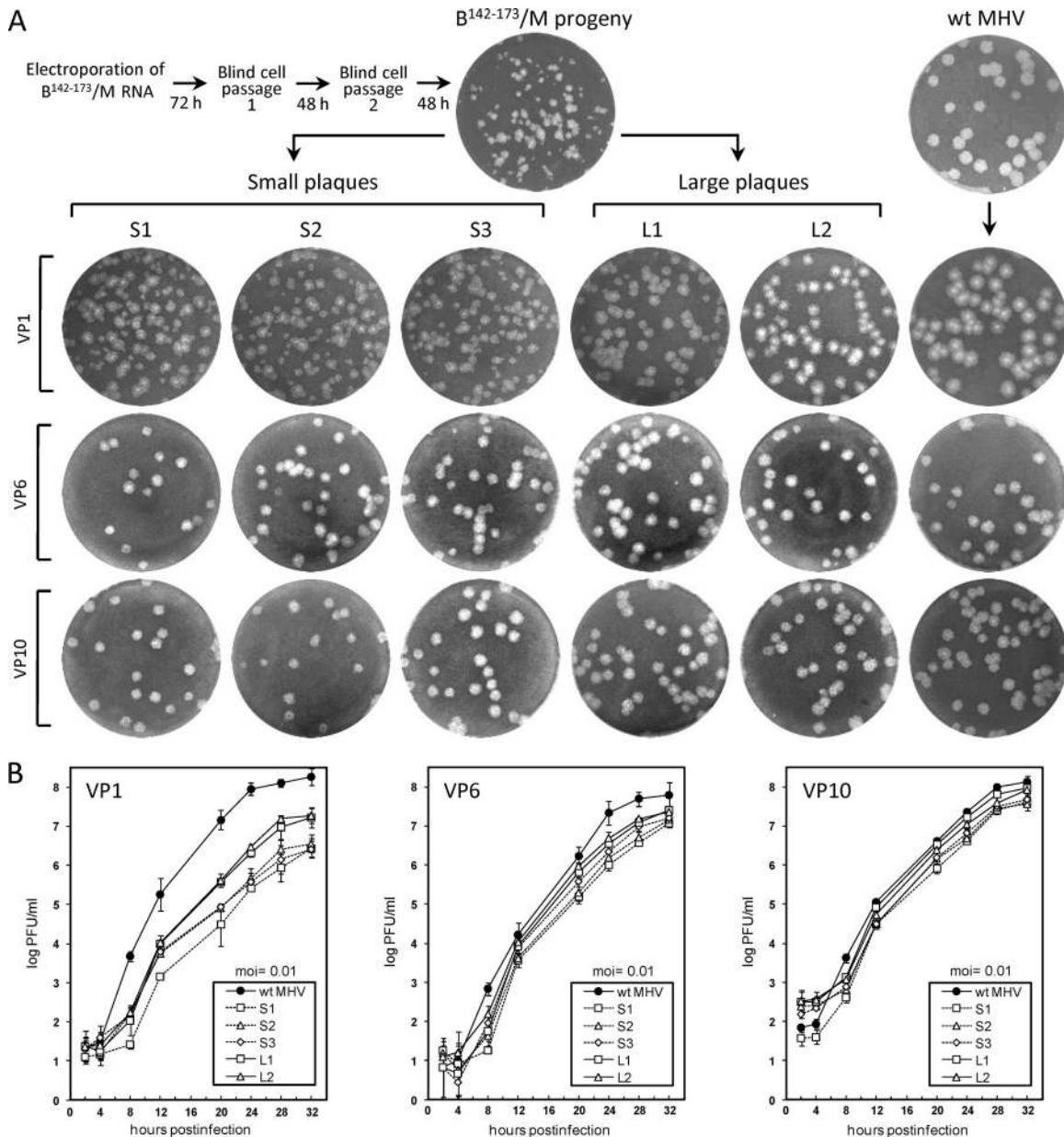
above, Mfold patterns that might identify base pairings for the long-range RNA-RNA interaction were sought by folding nt 1 to 400 or nt 1 to 500 of additional fully sequenced MHV-like and BCoV-like betacoronaviruses. Interestingly, base pairing between the inter-stem-loop domains (Fig. 4A) and respective partnering domains were readily found for the MHV-like, HCoV-HKU1, and BCoV-like coronaviruses (Fig. 4B) that revealed 25 to 27 potential base pairings within the 30- to 32-nt window. Figure 4B also depicts the phylogenetic covariation of sequences that maintain base pairings within the virus group. The Mfold and covariation results, therefore, appear to be consistent with a functional role for the base pairing as postulated above.

***In vitro* mutagenesis to form base pairing in the B<sup>142–173</sup>/M chimera with C158G, A163U, and G169U (thus recapitulating reverted virus S1 at passage 10) enabled immediate wt-like progeny after transfection of the chimeric genome.** To directly test the hypothesis that base-pairing between the inter-stem-loop and the Nsp1 partnering domains imparts fitness to the debilitated chimera, C158G, A163U, and G169U, the three putative suppressor mutations in isolate S1 at passage 10 (Table 1), were used to prepare chimera B<sup>142–173,S1</sup>/M (Fig. 5A), which was then tested by transfection. Whereas B<sup>142–173</sup>/M required two blind cell passages before the appearance of CPE (Fig. 2A), B<sup>142–173,S1</sup>/M developed CPE within 24 h posttransfection (hpt). Moreover, progeny from VP0 virus developed large plaques (Fig. 5B), had no additional putative suppressor mutations within the sequenced regions, had growth kinetics similar to that of wt MHV but with a 10-fold-lower final titer ( $10^7$  PFU/ml versus  $10^8$  PFU/ml for wt MHV) (Fig. 5C), and had a wt-like sgRNA profile as determined by Northern analysis (Fig. 5D).

These results, therefore, are consistent with the hypothesis that base pairing between the inter-stem-loop domain and the Nsp1 coding domain contributes to the replicating ability and fitness of MHV in cell culture.

**Preplaced suppressor mutations within the Nsp1 coding region of the otherwise B<sup>142–173</sup>/M chimeric RNA hastened the appearance of phenotypic revertants and suppressor mutations within the 5'-UTR-mapping inter-stem-loop domain.** To determine whether the rarer putative suppressor mutations within the Nsp1 coding region are sufficient for phenotypic reversion, the mutations A353U and A363G were separately placed into the MHV genome along with the wt BCoV inter-stem-loop domain and tested. With B<sup>142–173,A353U</sup>/M RNA, CPE appeared within 48 hpt and VP0 virus generated wt-like plaques, retained the A353U mutation, and also contained G169U within the transplanted domain (data not shown). With B<sup>142–173,A363G</sup>/M, CPE appeared within 48 hpt and VP0 virus generated wt-like plaques, retained the A363G mutation, and also contained G169U within the transplanted domain (data not shown). These results show that the A353U and A363G mutations each alone may not suffice as suppressor mutations but likely hastened the appearance of additional suppressor mutations within the transplanted pyrimidine-rich domain.

With the exception of the of a C30U mutation, single potential suppressor mutations occurring outside the long-range base-pairing windows, namely, C39U, U240C, and A591G, were not tested as suppressors (Table 1). When the C30U mutation alone was transplanted along with the BCoV inter-stem-loop domain into the MHV background, viable virus appeared, but so did C145 and G169 within the window. This result suggested that C30U may not



**FIG 2** Phenotypes of representative plaque-purified progeny from cells electroporated with chimeric  $B^{142-173}/M$  RNA. (A) Plaque appearances after culturing for 60 h on L2 cells. Isolated small plaques (S1 to S3 out of 6 total) and large plaques (L1 and L2 out of 11 total) were serially passed 10 times on DBT cells. (B) Growth kinetics for three isolated small plaques (hatched lines) and two isolated large plaques (solid lines) following their selection at viral passages 1, 6, and 10.

alone function as a suppressor but may by some unknown mechanism hasten the appearance of suppressor mutations. This phenomenon was not explored further.

**The BCoV 5' UTR and the (nearly) entire BCoV Nsp1 coding region function together in the MHV background to yield wt MHV-like progeny in cell culture.** The evidence showing that a virus species-specific interaction between the inter-stem-loop domain in the 5' UTR and its partnering domain in the Nsp1 coding region is required for virus replication led us to directly test a bidirectional matching of the two domains. Since it had been shown previously that the functional BCoV SLIV alone in the MHV genome extends 16 nt into the Nsp1 coding region (22), the

full BCoV 5' UTR in addition to the 16-nt extension (i.e., nt 1 to 226) was used for testing with the remaining MHV Nsp1 coding region in the MHV background. For this,  $B^{1-226}/M$  (Fig. 6A) was tested but was found to be nonviable over three blind cell passages from three separate electroporation attempts (Fig. 6B) (22), probably because of the incompatibility between the inter-stem-loop domain (from BCoV) and the Nsp1 partnering domain (from MHV) described above. Likewise, no viable progeny were generated when a chimera was tested in which the entire MHV 5' UTR plus its 16-nt extension into its Nsp1 coding region was used with the adapted BCoV Nsp1 coding region (i.e., one lacking the first 16 nt and last 42 nt) in the MHV background to form  $B^{227-906}/M$  (Fig.

TABLE 1 Potential compensatory mutations within the 5' UTR and Nsp1 coding region of progeny from the B<sup>142-173</sup>/M chimera

Clone <sup>a</sup>	Viral passage	Nt changes within 5' UTR							Nt changes within nsp1 cistron							Basepairs <sup>b</sup>	Folding <sup>c</sup>	
		C30	C39	C145	C158	U159	U160	A163	G169	U240	A347	C348	A353	A358	A363			A591
S1	VP1	U															22	
	VP6	U															23	
	VP10	U			G			U	U								24	1
S2	VP1	U															22	
	VP6	U		U													23	
	VP10	U		U													23	1-3
S3	VP1	U															22	
	VP6	U			G			U							G <sup>d</sup>		24	
	VP10	U			G			U	U						G <sup>d</sup>		25	1
S4	VP1	U										A <sup>e</sup>					23	
	VP6	U						U				A <sup>e</sup>					23	
	VP10	U		U				U				A <sup>e</sup>					24	2
S5	VP1	U				G											22	
	VP6	U		U		G		U	U								25	
	VP10	U		U		G		U	U								25	2
S6	VP1	U															22	
	VP6	U				G		U	U								24	
	VP10	U				G		U	U								24	2
L1	VP1	U			G												22	
	VP6	U			G			U									23	
	VP10	U			G			U									23	1
L2	VP1	U						U									22	
	VP6	U						U				U <sup>f</sup>					23	
	VP10	U						U				U <sup>f</sup>					23	1-3
L3	VP1	U						C									22	
	VP6	U						C								G <sup>g</sup>	22	
	VP10	U		U				C							G <sup>g</sup>		23	1-3
L4	VP1	U				G											22	
	VP6	U				G		U									23	
	VP10	U				G		U									23	2
L5	VP1	U						U									22	
	VP6	U						U						C <sup>h</sup>			23	
	VP10	U						U						C <sup>h</sup>			23	1-3
L6	VP1	U						U									23	
	VP6	U			U			U	U								24	
	VP10	U			U			U	U								24	1-3
L7	VP1	U		U				U									23	
	VP6	U		U				U									23	
	VP10	U		U				U									23	1-3
L8	VP1	U						U									23	
	VP6	U						U									23	
	VP10	U						U									23	1-3
L9	VP1	U										G					23	
	VP6	U						C				G					24	
	VP10	U						C				G					24	3
L10	VP1	U		U								C <sup>i</sup>					22	
	VP6	U		U		G											24	
	VP10	U		U		G											24	1
L11	VP1	U						U									23	
	VP6	U						U									23	
	VP10	U						U									23	1-3

<sup>a</sup> Plaque purified clones were selected at virus passage 0 (VP0) and identified as small (S) or large (L).

<sup>b</sup> Number of base pairs within the base-pairing window (identified as positions 145 to 169 in the 5' UTR and 347 to 363 in the Nsp1 cistron).

<sup>c</sup> See Fig. 3 for descriptions of the folding patterns.

<sup>d</sup> Causes K52E.

<sup>e</sup> Causes P47T.

<sup>f</sup> Causes K48N.

<sup>g</sup> Causes M128V.

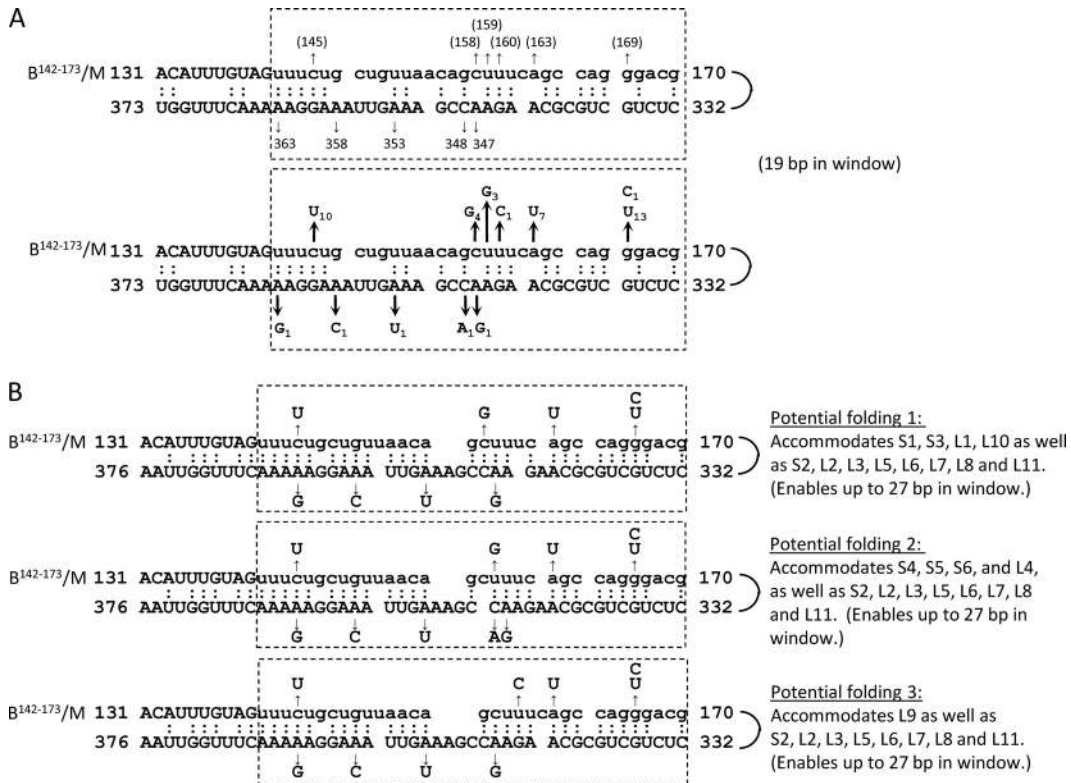
<sup>h</sup> Causes K50T.

<sup>i</sup> Causes F11L.

6A and B). Note that for this construct, the MHV 3' 42 nt of the Nsp1 coding region were retained to keep the region compatible with the MHV papain-like protease that separates Nsp1 and Nsp2 (17). However, when a construct in which the BCoV 5' UTR containing the 30-nt MHV inter-stem-loop domain was tested in the MHV background as in B<sup>1-141,174-226</sup>/M (Fig. 6A), CPE appeared within 48 hpt, and progeny from electroporated RNA (VP0) yielded medium plaques, a peak titer of  $1.0 \times 10^6$  PFU/ml, and a full set of viral sgRNAs (Fig. 6B to D). These results together demonstrated a requirement for compatibility between the 5'-UTR-mapping inter-stem-loop domain and its cognate partnering domain within the Nsp1 coding region. Conversely, when a construct with the entire MHV 5' UTR but with the BCoV 32-nt inter-stem-loop domain and the BCoV Nsp1 coding region in the MHV background, B<sup>142-173,227-906</sup>/M (Fig. 6A), was tested, CPE was observed within 48 hpt, and viable progeny (VP0) producing large plaques, a titer of  $1.0 \times 10^6$  PFU/ml, and a full set of viral sgRNAs (Fig. 6B to D) were obtained. This construct, therefore, also demonstrated a compatibility requirement between the 5' UTR-mapping inter-stem-loop domain and its cognate partnering domain within the Nsp1 coding region.

These results are fully consistent with the notion that functional base pairing between the two domains is needed for virus replication. Still, the compatibility in these constructs was not perfect, since the titer of progeny was 100-fold lower than that for wt MHV. Interestingly, when the entire BCoV 5'-proximal region of the genome through the Nsp1 coding region to nt 906 was tested in construct B<sup>1-906</sup>/M, CPE was found within 24 hpt and viral progeny produced plaques, growth kinetics, a final titer, and a sgRNA profile that were indistinguishable from those of wt MHV (Fig. 6A to D). These results further demonstrate a requirement for compatibility between the 5' UTR and cognate Nsp1 coding region for BCoV and MHV when used in the MHV background for replication and fitness studies in cell culture. Without a reverse genetics system for BCoV, we could not apply the reciprocal test for this finding.

**The 30-nt inter-stem-loop domain in the 5' UTR is not an absolute requirement for MHV replication in cell culture.** Since in MHV, distinct regions in the 5' UTR, including SLIII and the loop region of SLIV (Fig. 1), and a large region of the 3' UTR have been shown to be dispensable for virus replication in cell culture (20, 22, 50), this test was applied to the 30-nt inter-stem-loop domain in MHV. Surprisingly, a deletion of nt 141 to 170 in the MHV genome (Fig. 7A) was not lethal, but rather, CPE appeared after one blind cell passage following transfection of the mutated genomic transcript, and progeny showing wt-like large plaques and a final growth titer of  $1.0 \times 10^6$  PFU/ml were obtained (Fig. 7B and C). Interestingly, an A649C mutation (H147P) was found in the Nsp1 coding region of all progeny, and G516A (V103I), C583U (T125I), and A880C (E224A) mutations were found in some but not all progeny at VP10 (Fig. 7C). Furthermore, a UCUAA sequence was found inserted within the upstream transcription-regulatory sequence of all progeny by VP10 (Fig. 7A and C). These results together with the observation of very poor viral growth when the BCoV inter-stem-loop domain replaced the homologous native MHV domain demonstrate that the inter-stem-loop domain is required for optimal viral fitness but is not absolutely required for viral replication in cell culture.



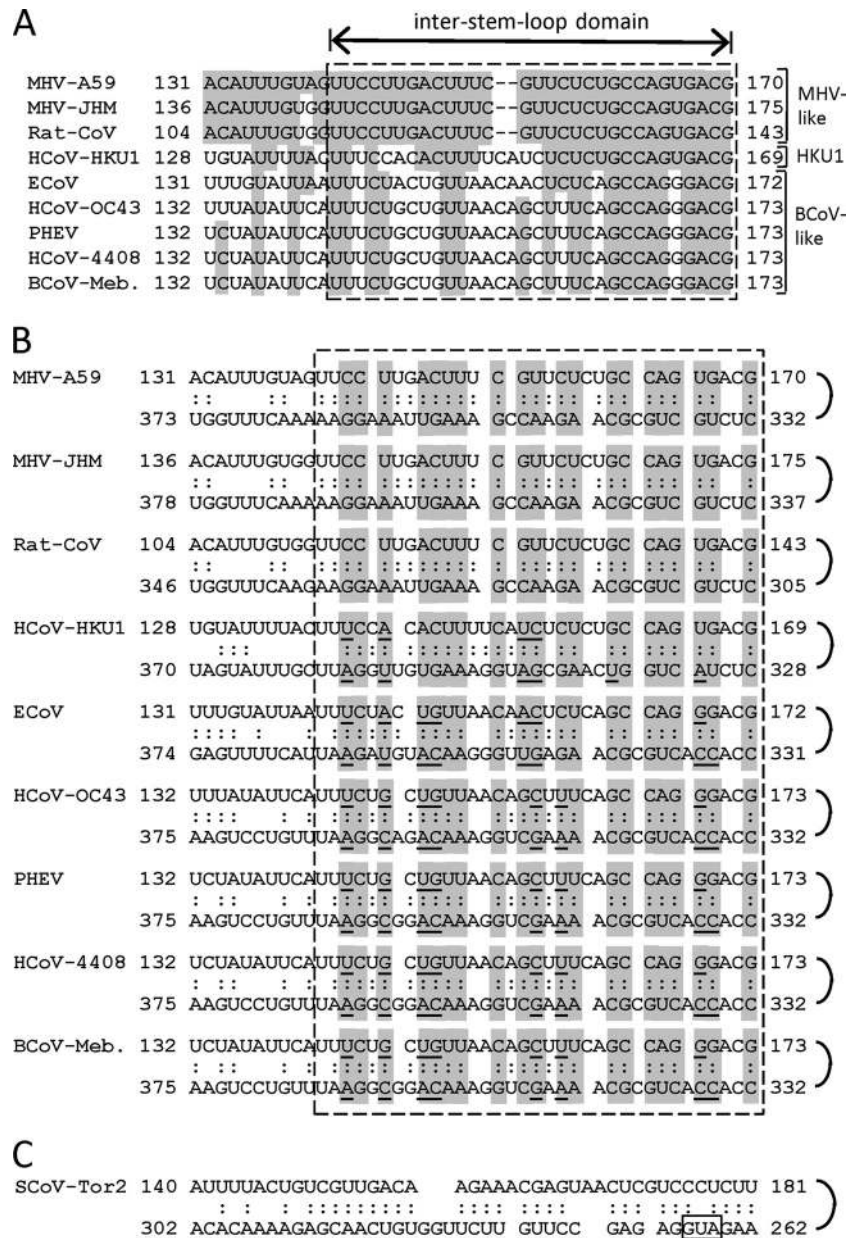
**FIG 3** Potential suppressor mutation-induced base pairings in progeny of the B<sup>142-173</sup>/M chimera. (A) Summary of potential suppressor mutations and their positions within the inter-stem-loop and downstream partnering domains that appeared in the small and large plaque isolates after 10 serial passages (depicted as in Fig. 1C). (Top) Base positions (in parentheses) for the BCoV sequence (lowercase) use BCoV numbering. Base positions elsewhere use MHV numbering. (Bottom) Summary of base changes at the named positions. The subscript numbers indicate the number of occurrences in 17 total isolates, shown in Table 1. (B) Three potential folding patterns as suggested by Mfold that together maximize the base pairings enabled by the suppressor mutations. Any of the three potential foldings can accommodate revertants S2, L2, L3, L5, L6, L7, L8, and L11, and others are accommodated as indicated. For revertants depicted in Table 1, base pairings within the base-pairing window at passage 10 increased from 19 to 22, 23, 24, or 25 (as summarized in the last two columns of Table 1).

**DISCUSSION**

Here, we report the discovery that for the bovine coronavirus and mouse hepatitis coronavirus, separate species in the (formerly group 2a) betacoronavirus genus, there exists a functional interdependence between their respective 5' UTRs and Nsp1 coding regions for optimal virus growth in cell culture and that part of that interdependence is a long-range (~200-nt) RNA-RNA interaction. Furthermore, it was observed that whereas only some parts of the BCoV 5' UTR and Nsp1 coding region can be transplanted into the MHV genome to directly yield viable virus and some parts cannot (22), when the entire BCoV 5' UTR and Nsp1 coding region are transplanted together as a cognate unit, nearly fully fit MHV wt-like progeny are directly obtained, and there appear to be no interspecies constraints on replication. These results therefore show that the BCoV *cis*-acting RNA elements in the 5' UTR and Nsp1 coding region, although evolutionarily divergent (~36% nucleotide divergence in the 5' UTR, ~40% nucleotide divergence in the Nsp1 coding region, and ~53% amino acid divergence in the Nsp1 protein), are successfully recognized as a cognate unit and utilized by the MHV proteins for genome replication and packaging with apparently no detrimental effect. The long-range RNA-RNA interaction function for replication also explains, at least in part, the previously reported *cis*-replication requirement for the 5'-terminal part of the Nsp1 coding region in BCoV defective interfering (DI) RNA (7, 8) and the presence of

this region in all known naturally occurring group 1 and group 2 coronavirus DI RNAs (3, 4). They possibly also explain, in part, the requirement for this region in the MHV genome for virus replication (5).

Is the identified long-range RNA-RNA interaction a common feature of coronaviruses? In a recent comprehensive prediction of coronavirus 5'-end-proximal higher-order structures based predominantly on Mfold and phylogenetic and covariation analyses by Chen and Olsthoorn (10), seven substantial stem-loops (1 through 5, and unnamed 6 and 7) were described for the 5'-terminal 340 nt of group 2a coronaviruses that, for BCoV, are essentially identical to the seven stem-loops depicted in Fig. 1B (SL1, -2, -II, -III, and -IV) and in references 23 (SLV) and 7 (SLVI) (sequence differences are due to the different strains of BCoV used, BCoV-Mebus versus BCoV-Ent). It is of interest that for both BCoV and MHV, stem-loops V and VI (equivalent to stem-loops 6 and 7 in reference 10; also data not shown) map entirely within the Nsp1 coding region and (except for possibly a short region as noted below) lie between the upstream and downstream partnering domains in the long-range RNA-RNA interaction. A possibly analogous long-range RNA-RNA interaction is predicted for each of the group 2b, 2c, and 2d coronaviruses as well, but in these cases, in contrast to the group 2a coronaviruses, the long-distance upstream and downstream base-pairing partners are contained



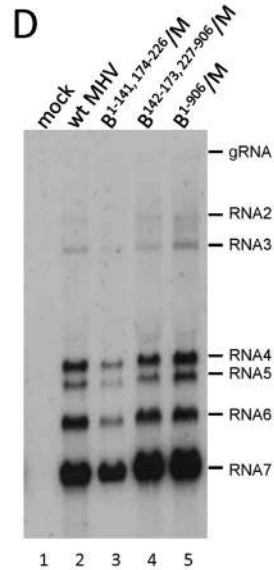
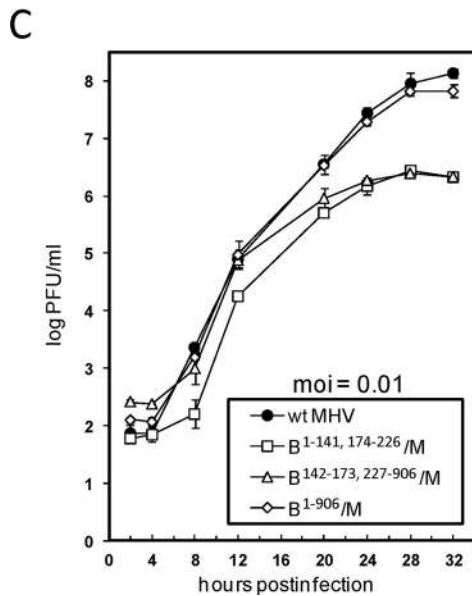
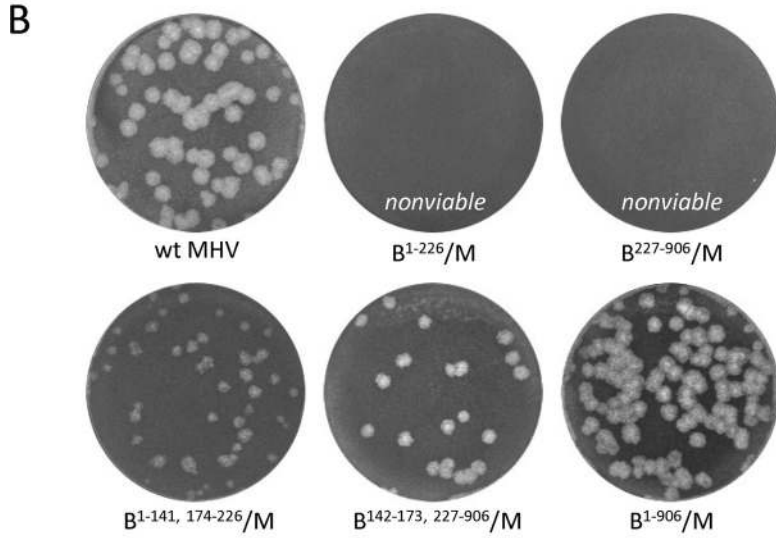
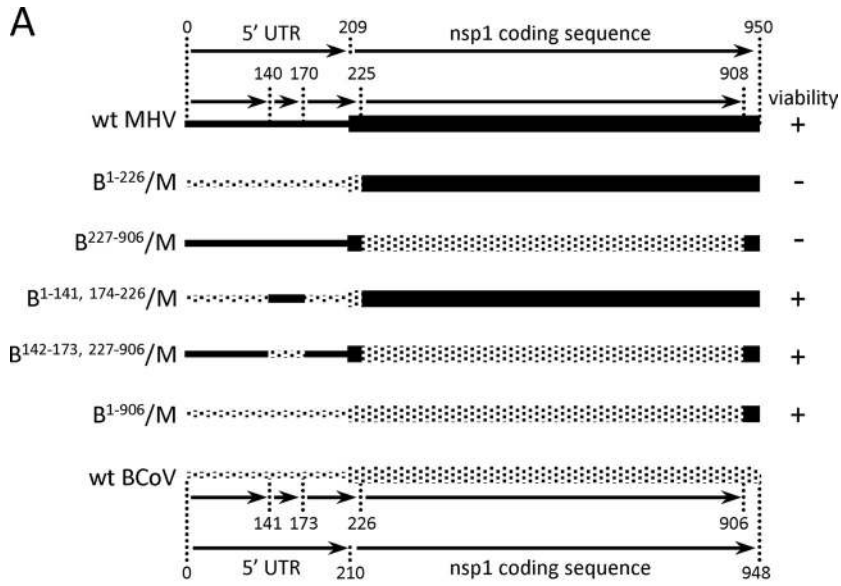
**FIG 4** Base pairings between the inter-stem-loop and partnering domains for 10 betacoronaviruses. (A) Alignment of the inter-stem-loop domains for MHV-like, HCoV-HKU1, and BCoV-like betacoronaviruses. Shading identifies bases that are in common with MHV-A59. The inter-stem-loop domains are identified with a hatched box. (B) Base pairings as predicted by Mfold. Shaded areas identify regions of base pairing common to the whole group. Underlined bases within the shaded areas are variant bases relative to those in MHV-A59 that maintain base pairing at the same position. Note that the number of base pairings within the inter-stem-loop domain for any given virus ranges from 25 to 27. (C) Potential 5'-end-proximal long-range RNA-RNA interaction derived by Mfold analysis of the 5' region of the SCoV-Tor2 genome. This folding prediction is identical to that named for stem-loop 5 by Chen and Olsthoorn (10). The AUG start codon for the Nsp1 coding region is identified with a hatched box.

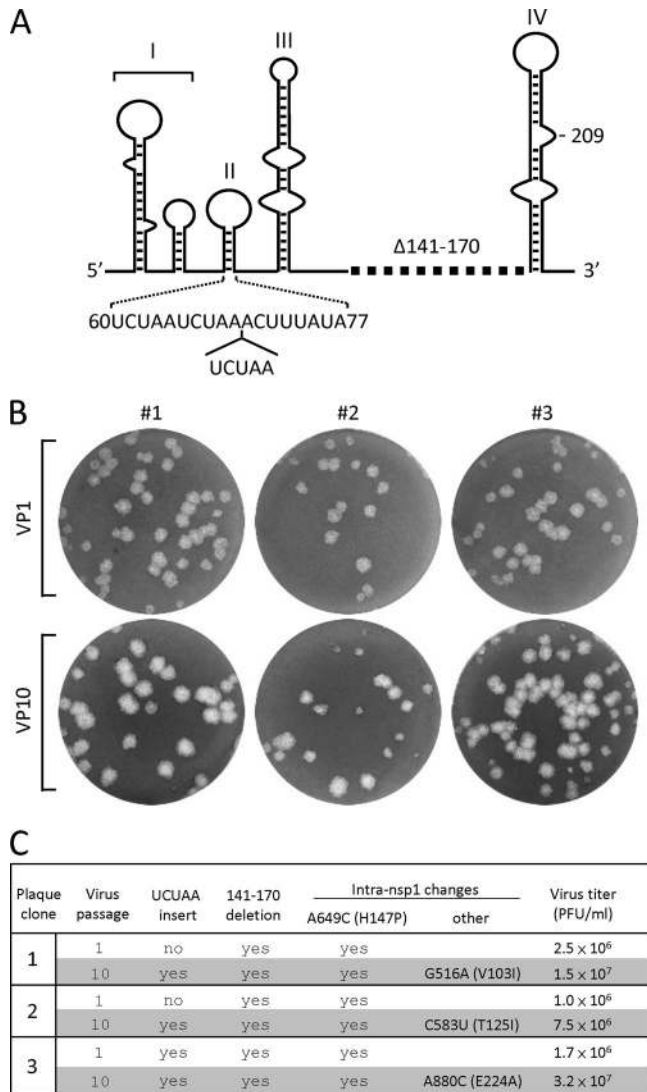
entirely within the stem of SL5 (equivalent to SLIV in Fig. 1; see also Fig. 4C, and see Fig. 3B in reference 10). SL5 for groups 2b, 2c, and 2d coronaviruses also harbors a large terminal loop that, in turn, contains three smaller stem-loops with primarily hexameric (UUYCGU) terminal loops (10). That is, for the group 2b SCoV-like coronaviruses (harboring an 80-nt terminal loop), the group 2c BCoV-HKU5-1-like coronaviruses (harboring a 94-nt terminal loop), and the group 2d BCoV-HKU9-1-like coronaviruses (harboring a 55-nt terminal loop), there is a long-

range RNA-RNA interaction, but the upstream portion is nearly entirely within the 5' UTR and the downstream portion is confined to the 5'-terminal 29 to 113 nt of the Nsp1 coding region. In a pattern similar to that of the group 2b through 2d coronaviruses, long-range RNA-RNA interactions that might also be analogous are predicted within SL5 of the group 1a TGEV-Purdue-like coronaviruses (with a 106-nt terminal loop), the group 1b HCoV-229E-inf-1-like coronaviruses (with a 100-nt terminal loop), the group 1c PEDV-CV777-like coronaviruses (with a 116-nt termi-









**FIG 7** Deletion of the 30-nt inter-stem-loop domain in MHV leads to viable but debilitated progeny. (A) Schematic depiction of the MHV 5' UTR with a deleted inter-stem-loop region ( $\Delta 141-170$ ) and the inserted UCUAA element. The UCUAA insertion could theoretically have occurred following nt 59, 64, or 69 (illustrated as the latter). (B) Plaque patterns for plaque clones 1 to 3 at virus passages 1 and 10. (C) Genotypes and virus titers for the viruses depicted in panel B.

long-range RNA-RNA interaction, however, we noted that the downstream 9 nt of SLVI in BCoV (nt 332 to 340) (7) are also part of the downstream base-pairing window in the long-range RNA-RNA interaction (Fig. 1B). That is, the mutations that destabilized the lower stem of SLVI in BCoV DI RNA (C333G and C336U) (7) would also have decreased base pairing in the long-range RNA-RNA interaction from 25 to 24 (Fig. 1B).

Thus, either disruption (i.e., that in SLVI or that in the long-range RNA-RNA interaction) may have been the lesion that blocked DI RNA replication. This arrangement may suggest a possible functional switching between two alternate structures. A similar arrangement involving the four 3'-terminal nucleotides in MHV SLVI (nt 332 to 335) is also possible (data not shown).

Inasmuch as the 5' UTR-mapping 30-nt inter-stem-loop component of the long-range RNA-RNA interaction can be deleted without killing MHV replication (Fig. 7), the question of what role(s) the long-range RNA-RNA interaction might play that could be considered optional for virus survival arises. Currently, we can only speculate from known functions of 5'-end-proximal *cis*-acting replication elements in coronaviruses and other plus-strand RNA viruses (see the introduction). Based on available data, it is plausible that the long-range RNA-RNA interaction plays a role in one or both of two potential regulatory pathways. First, it might regulate an optional non-canonical translation initiation site in a manner that ensures viral genome translation under conditions of cell stress when canonical translation initiation mechanisms are largely repressed (1, 26, 30, 44). This role is suggested by the demonstration of a potential internal ribosomal entry site on group 3 coronavirus genomic RNA that needs yet to be demonstrated in other coronavirus groups (34) and by robust translation of MHV RNAs during global inhibition of cellular mRNA translation (2, 40, 45). Such an alternate mechanism of translation initiation has been shown to function on some capped and polyadenylated viral and cellular mRNAs, to be influenced by double-stranded RNA regions near the translation start codon, and to be under the control of *trans*-acting proteins (1, 26, 30, 44).

A second potential regulatory pathway for the long-range RNA-RNA interaction could be maintenance of a proper acceptor orientation for the RNA-dependent RNA polymerase (RdRp) template switching during generation of minus-strand templates for subgenomic mRNA synthesis (43, 49, 55). This is suggested by the appearance of the UCUAA component within the UCUAAAC intergenic transcriptional regulatory sequence following deletion of the 30-nt inter-stem-loop region (Fig. 7). Addition of the UCUAA element in the same region was also found when stem-loop 4 (referred to as SLIII in Fig. 1A) was deleted, which led to the suggestion that stem-loop 4 functions as a required spacer element for sgRNA synthesis (50). In earlier studies, the UCUAA element was also found to spontaneously appear at this site in some MHV strains in which its occurrence positively correlated with enhanced transcription (37, 51, 53). Regulation of mRNA synthesis is also suggested by the increased synthesis of sgRNAs when amino acid-changing mutations were made in the (then unknown) downstream partnering domain of the MHV long-range interaction during the analysis of Nsp1 function by Brockway and Denison (5). For Nsp1 functional analysis in MHV-A59, 13 amino

**FIG 6** Requirement for compatibility between the 5'-UTR-located inter-stem-loop and downstream partnering domains for MHV replication. (A) For viability, the chimeric virus must contain the MHV inter-stem-loop domain within the BCoV 5' UTR when in the MHV background (as in B<sup>1-141,174-226</sup>/M), the BCoV inter-stem-loop domain and BCoV Nsp1 coding region when in the MHV background (as in B<sup>142-173,227-906</sup>/M), or the BCoV 5' UTR and BCoV Nsp1 coding region when in the MHV background (as in B<sup>1-906</sup>/M). Note that the in-frame C-terminal fusion site for the Nsp1 coding region occurs between BCoV nt 906 and MHV nt 909. (B) Plaque phenotypes for the progeny from the variant constructs described for panel A. (C) Growth kinetics of the progeny depicted in (B). (D) Northern analyses of the progeny depicted in for panel B.

acids within the amino-terminal third of Nsp1 were changed to alanine by reverse genetics and tested for viability. Although only some of the mutations were found to be lethal, two tested simultaneously, E46A and K48A, made by A346C, A351G, and A352C mutations, were not lethal but inexplicably caused an ~20% increase in overall viral RNA synthesis (5). Although these three mutations which map within the long-range base-pairing window would have decreased the base pairings only within the window from 27 to 25, there might have been enough RNA rearrangement to affect upstream RdRp template switching and an increase in subgenomic mRNA synthesis. Alternatively, the amino acid changes could have caused alternate template switching behavior by an entirely different unknown mechanism.

From the present study, there is no clear evidence to suggest that MHV behavior was altered as a function of amino acid changes arising from suppressor mutations. However, in no case was an amino acid-changing mutation found in the absence of other potential suppressor mutations, making it unfeasible to assess the effects of single amino acid changes alone. Of the five suppressor mutations within the downstream Nsp1 coding region, four (S3, S4, L2, and L5) led to amino acid changes within Nsp1 (K52E, P47T, K48N, and K50T, respectively) (Table 1), but progeny from each of these made large plaques (Fig. 2A and data not shown), and two, S3 and L2, had near-wt growth kinetics at passage 10 (Fig. 2B). No evaluation of viral RNA levels was made for any of these, however.

## ACKNOWLEDGMENTS

We thank Ralph Baric for the MHV-A59-1000 reverse genetics system.

This work was supported by Public Health Service grant AI014367 from the National Institute of Allergies and Infectious Diseases and by funds from the University of Tennessee Center of Excellence in Livestock Diseases and Human Health.

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