Downstream Ribosomal Entry for Translation of Coronavirus TGEV Gene 3b

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Gene 3b (ORF 3b) in porcine transmissible gastroenteritis coronavirus (TGEV) encodes a putative nonstructural polypeptide of 27.7 kDa with unknown function that during translation in vitro is capable of becoming a glycosylated integral membrane protein of 31 kDa. In the virulent Miller strain of TGEV, ORF 3b is 5'-terminal on mRNA 3–1 and is presumably translated following 5' cap-dependent ribosomal entry. For three other strains of TGEV, the virulent British FS772/70 and Taiwanese TFI and avirulent Purdue-116, mRNA species 3–1 is not made and ORF 3b is present as a non-overlapping second ORF on mRNA 3. ORF 3b begins at base 432 on mRNA 3 in Purdue strain. In vitro expression of ORF 3b from Purdue mRNA 3-like transcripts did not fully conform to a predicted leaky scanning pattern, suggesting ribosomes might also be entering internally. With mRNA 3-like transcripts modified to carry large ORFs upstream of ORF 3a, it was demonstrated that ribosomes can reach ORF 3b by entering at a distant downstream site in a manner resembling ribosomal shunting. Deletion analysis failed to identify a postulated internal ribosomal entry structure (IRES) within ORF 3a. The results indicate that an internal entry mechanism, possibly in conjunction with leaky scanning, is used for the expression of ORF 3b from TGEV mRNA 3. One possible consequence of this feature is that ORF 3b might also be expressed from mRNAs 1 and 2.

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Key Words: porcine transmissible gastroenteritis coronavirus; gene 3b; ribosomal scanning; ribosomal shunting.

INTRODUCTION

Expression of coronavirus genes occurs through the synthesis of a 3' coterminal nested set of mRNAs. Although coronavirus mRNAs are structurally polycistronic (the 3'-most mRNA in many but not all viral species is monocistronic), evidence from studies of translation both in vitro and in vivo has suggested that most function as monocistronic messages. That is, despite their polycistronic configuration, usually only the 5'-terminal ORF on each is abundantly translated (reviewed in Lai and Cavanagh, 1997, and Luytjes, 1995). The location of some coronavirus genes, however, is not 5'-terminal on any mRNA, which would require that the gene, if expressed, be translated by a mechanism allowing translation reinitiation, leaky scanning, frameshifting, or a downstream entry of ribosomes. Examples of such genes include (1) ORF 1b on mRNA 1, from which a polyprotein is synthesized following 1 ribosomal frameshifting (Brierly et al., 1987; Eleouet et al., 1985); (2) ORFs 3b and 3c on avian infectious bronchitis virus mRNA 3 (Bournsell et al., 1985), from which 7.4- and 12.4-kDa proteins are synthesized following leaky scanning and internal ribosomal entry, respectively (Liu et al., 1991; Liu and Inglis, 1992; Le et al., 1995); (3) ORF 5b on mouse hepatitis virus mRNA 5 (Skinner et al., 1985), from which a 9.6-kDa protein (the E protein) is synthesized (Budzilowicz and Weiss, 1987; Leibowitz et al., 1988), by an apparent internal ribosomal entry mechanism (Thiel and Siddell, 1994); and (4) the I ORF in mRNA 7 of the bovine and mouse hepatitis coronaviruses, from which the I protein is made in the +1 reading frame relative to N (Senanayake et al., 1992; Fischer et al., 1997) following ribosomal scanning (Senanayake and Brian, 1997).

In this study, we examine the mechanism by which gene 3b is expressed from mRNA 3 in the Purdue strain of TGEV and demonstrate that, surprisingly, it may be approached by ribosomes entering internally and not necessarily through a leaky scanning step as would be predicted from mRNA 3 sequence. Gene 3b in TGEV is unusual in that for one strain of virus, the virulent Miller strain (Wesley et al., 1989), it is expressed as the 5'-terminal ORF on mRNA 3–1, whereas in three other strains, the virulent British FS772/70 and Taiwanese TFI strains and the avirulent Purdue-116 strain, it is expressed as the second ORF on mRNA 3 (Britton et al., 1989; Chen et al., 1995; this study) (note mRNA structures in Fig. 1A). The differences in transcription patterns appear to be a function of the canonical TGEV UCIAAAC intergenic sequence positioned 18 nt upstream of gene 3b in the genome, which in the Miller virus totally conforms to the canonical sequence but in the British FS772/70 and Purdue-116 strains is UCIAAAA and in the Taiwanese TFI strain is ACIAAAC. The nonconforming intergenic sequences apparently fail to promote syne-
FIG. 1. TGEV genome map and the cDNA constructs used for the expression of gene 3b as a downstream ORF. (A) TGEV genome map showing the positions of cDNA clones FT44 and FT39 from which the expression clones were derived, and the gene maps of mRNA species 1, 3, and 3–1. (B) Map positions of the start and stop codons for ORFs 3a, 3b, and 4 in mRNA 3. (C) cDNA constructs used for the synthesis of RNA transcripts from which translation of ORF 3b was studied.
sis of a subgenomic mRNA. Translation from ORF 3b when it occurs as the second ORF on mRNA 3 must, therefore, require either a reinitiation of translation after translation of the upstream ORF, a leaky scanning by ribosomes over a long distance (431 nt; Fig. 1B), or a downstream entry of ribosomes.

Because gene 3b is not intact in some strains of TGEV (i.e., it is either severely truncated by frameshift mutations as in the Purdue-115 strain [Rasschaert et al., 1987] or by deletions as in the avirulent Miller strain [Wesley et al., 1990, 1991]) and cannot produce a full-length product, it has been suggested that its product fulfills a specialized function, perhaps during animal infection (reviewed in Enjuanes et al., 1995), and is not required for virus replication. Similar conclusions were reached after revelations of a truncated gene 3b in strains of the closely related porcine epidemic diarrhea virus (Vaughn et al., 1995). It might therefore be assumed that gene 3b is not translated when it occurs as a downstream ORF as in mRNA 3. However, a product from gene 3b is made in cells infected with the Purdue-116 strain of virus (O’Connor and Brian, 1999), indicating a mechanism must exist for its synthesis from mRNA 3. Here we report that, whereas mRNA 3 has a sequence predicting leaky scanning for the translation of ORF 3b by the model of Kozak (1989), experimental results with mutant constructs suggested downstream entry of ribosomes might also be used. Furthermore, deletion analysis indicated that the internal entry of ribosomes did not depend on an immediate upstream internal ribosomal entry structure (IRES) and suggested ribosomes are entering very close to the ORF 3b start site by a mechanism resembling shunting.

RESULTS

mRNA 3, but not mRNA 3–1, is made in cells infected with the Purdue-116 strain of TGEV

Northern analyses of Purdue-116 virus-infected cells carried out previously in our laboratory with oligonucleotide probes specific for the 3′ end of the genome (i.e., a sequence from within the 3′-proximal HP ORF) had identified eight species of mRNA, leading us to conclude that ORFs encoding the 7.7-kDa (gene 3a) and 27.7-kDa (gene 3b) proteins are each 5′-terminal on separate mRNA species (Sethna et al., 1989). To test this conclusion, separate Northern analyses were done with probes specific for the 90-nt leader and for genes 3a and 3b. Our rationale was that mRNA species 3 and 3–1 would be distinguishable with probes binding within ORFs 3a and 3b because transcripts of 3875 and 3561 nt (or even 4075 and 3761 nt if they included poly A tails of 200 nt in length) are resolvable on a gel of 1% agarose. Northern analyses with the separate probes revealed bands with identical mobilities, indicating the presence of mRNA 3, but not mRNA 3–1, in RNA from Purdue-116 virus-infected cells (Fig. 2A, lanes 4–9).

To test this conclusion by a second method, RT-PCR analysis was done with oligonucleotide primers specific to gene 3b and the minus strand of the leader. Amplified products of 463 and 876 nt would be expected from mRNAs 3–1 and 3, respectively. A product of 5248 nt
might also be found from mRNA 2, the mRNA encoding the spike protein. From RT-PCR analysis only a single product of 876 nt with the proper sequence as determined by cloning and sequencing was obtained (Fig. 2B, lane 3; sequencing data not shown), indicating the presence of mRNA 3 but not mRNA 3–1. To establish that the experimental protocol would have detected mRNA 3–1 if present, RNA was extracted from Miller virus-infected cells and used in parallel. From this, the expected 463-nt mRNA 3–1-derived product was obtained (Fig. 2B, lane 4). No product was obtained with RNA from uninfected cells (Fig. 2B, lane 2).

Genes 3a and 3b, but not gene 4, are translated in vitro from synthetic mRNA 3-like transcripts containing all three ORFs

To test by in vitro translation whether the 27.7- and 20-kDa gene 3b products (O’Connor and Brian, 1999) are synthesized when ORF 3b is positioned downstream of ORF 3a (beginning at base 337) on synthetic transcripts, uncapped transcripts of pORF3a-3b-4 DNA linearized at the BamHI site 50 nt downstream from the stop codon of gene 4 (Fig. 1C) were translated in either wheat germ extract or rabbit reticulocyte lysate. In both, products from genes 3a (the 7.7-kDa protein containing one methionine) and 3b (the 27.7-kDa form of the protein containing eight methionines and the 20-kDa form presumably containing seven methionines), but not gene 4 (the 9.2-kDa E protein containing four methionines), were obtained (results for wheat germ extract are shown in Fig. 3A, lane 5; note the marker positions in lanes 2–4 and the absence of endogenous product in lane 1). Because a nearly 27-kDa protein was also synthesized from an endogenous transcript in (some) rabbit reticulocyte lysates and the presence of abundant globin protein in the lysate interfered with the resolution of small proteins (data not shown), all subsequent studies described were carried out in wheat germ extract. Thus, from uncapped transcripts bearing similarity to mRNA 3 the first methionine codon in gene 3b (for synthesis of the 27.7-kDa protein) and also the second (assuming synthesis of the 20-kDa protein initiates at the second methionine codon [O’Connor and Brian, 1999]) are accessed for translation. On a molar basis, the amount of 27.7-kDa gene 3b product is approximately one-fifth of that from gene 3a (Fig. 3B). There was no evidence of a 9.2-kDa E protein from ORF 4 in this (Fig. 3A, lane 5) or in subsequent experiments, indicating ribosomal accessibility of ORF 3b was probably not the result of template fragmentation.

An upstream leader-containing sequence in transcripts of pLORF3a-3b-4 (Fig. 1C), although containing an additional 14 nts not found on mRNA 3, had only a small effect on the rate of translation from ORF 3b relative to 3a (Fig. 3A, lane 7, and Fig. 3C), indicating the leader sequence may not strongly influence translation from the downstream ORF; however, this needs confirmation with transcripts precisely mimicking the 5’ end of mRNA 3. As with transcripts of pORF3a-3b-4, no product was evident from ORF 4.

Translation of gene 3b from mRNA 3-like transcripts shows a pattern not fully consistent with a leaky scanning model

From precedents in eukaryotic mRNAs, it is unlikely that ribosomes would approach ORF 3b on mRNA 3 by a mechanism of translation reinitiation, since three in-frame strong stop codons follow ORF 3a (Fig. 1B). However, an approach by leaky scanning according to the model of Kozak (1989, 1991a,b) might be expected, since the initiator codon for ORF 3a (UGUA123UGG, in the
same reading frame as ORF 3b) and for three other potential small ORFs within gene 3a (UAGA167UGC, CAUA283UGC, and UCCA424UGC, all in the +1 reading frame relative to ORF 3b) are within contexts considered weak for initiation, whereas that for ORF 3b (AAA432UGA) is considered relatively strong. To test for ribosomal scanning on mRNA 3-like transcripts, three approaches were taken. In the first, the effect of a 5’ cap on the synthesis of 3a and 3b gene products was measured. Increased synthesis from both would be expected if 5’ cap-dependent entry followed by leaky scanning were used (Kozak, 1989, 1991a). Increased synthesis from 3b might also be expected if a cap-dependent shunting mechanism were used (Jackson, 1996; Mathews, 1996). As can be observed in Fig. 3A, lanes 6 and 8, and Figs. 3B and 3C, enhanced translation of both ORFs 3a and 3b resulted when capped transcripts of pORF3a-3b-4 and pLORF3a-3b-4 were translated. These results are therefore consistent with the mechanisms of leaky scanning and cap-enhanced ribosomal shunting.

In the second approach, the competitive effect of a soluble cap analog on the translation of ORFs 3a and 3b from capped transcripts of pORF3a-3b-4 was measured. With either leaky scanning or cap-dependent shunting, but not with cap-independent internal entry, competitive inhibition of translation from both ORFs would be expected (Iizuka et al., 1994; Jackson, 1996; Mathews, 1996). Nearly the same rate of inhibition was found, 70–75% with 0.1 mM and 85–90% with 2 mM cap analog (Figs. 4A and 4C), indicating either mechanism of cap-dependent entry could be functioning in the translation for ORF 3b.

In the third approach, the sequence context surrounding the 3a start codon in transcripts of pORF3a-3b-4 was modified to become strongly favorable for translation (GCCGCCATGG) (Kozak, 1991b) and the relative amounts of 3a and 3b gene products were measured. With leaky scanning, a diminished synthesis from 3b relative to 3a would be expected regardless of the capped status of the transcripts (Kozak, 1991b), whereas with shunting a change in the relative amounts would not necessarily be expected. As can be noted in Figs. 5A and 5B, whereas the accumulation of 3a product increased almost 20% relative to 3b with the improved Kozak consensus for capped transcripts, no increase was observed with uncapped transcripts. Intriguingly, the absolute amount of gene 3b product appeared nearly identical under all conditions of translation. These results, therefore, are not fully consistent with the leaky scanning model for ORF 3b.

FIG. 4. Competitive effects of soluble methylated cap analog on translation. (A) Results from pORF3a-3b-4. (B) Results from psCATORF3a-3b-4. Radiolabeled products from in vitro translation of synthetic transcripts were separated by SDS polyacrylamide gel electrophoresis and bands were quantitated by AMBIS radioanalytic imaging analysis of the dried gel. The quantitated results are shown for pORF3a-3b-4 (C) and for psCATORF3a-3b-4 (D). Note that only detected proteins are represented in the bar graph.
Translation and suggest the possibility of an internal entry of ribosomes.

Translation of ORF 3b is not blocked by the upstream insertion of a 884-nt-long sequence containing three sequential ORFs

To test for an internal entry of ribosomes onto ORF 3b, pORF3a-3b-4 was modified to psCATORF3a-3b-4 by the placement of an 884-nt-long sequence containing three sequential ORFs upstream of ORF 3a (Fig. 1C) and the products of translation were quantitated. An internal entry of ribosomes, either directed by an IRES element or by a shunting mechanism, would typically not be blocked by the presence of upstream ORFs of this dimension (reviewed in Jackson et al., 1995; Jackson, 1996; Mathews, 1996). Transcripts of psCATORF3a-3b-4 possessed a 5’ UTR of 59 nt; a five-methionine-containing 450-nt sCAT ORF beginning within an excellent Kozak context (AAAATGG) at base 60; a five-methionine-containing 4.4-kDa protein-encoding ORF beginning within a fair Kozak context (ATCATG) at base 346; a one-methionine-containing 4.5-kDa protein-encoding ORF beginning within a fair Kozak context (CCGAATG) at base 534; and ORF 3a beginning at base 944, ORF 3b beginning at base 1253, and ORF 4 beginning at base 1974. In addition, there is a 66-nt protein-encoding ORF beginning within a fair Kozak context (TCCATG) at base 346 within the sCAT ORF (in the +1 reading frame relative to sCAT).

These results show that translation of ORF 3b positioned 1263 nt downstream from the 5’ terminus in the synthetic construct is initiated by some form of internal entry of ribosomes and not by scanning, is influenced by a cap, and is not the result of a fragmented template.

Deletion analysis failed to identify a postulated IRES-like element within the upstream gene 3a

Although no universally identifying primary or secondary structural features of IRES elements are known, certain secondary structural features do appear necessary for IRES function (reviewed in Jackson, 1996). Within TGEV gene 3a, secondary structures can be predicted (Figs. 6A and 6B) that share features with the putative IRES element in IBV mRNA 3 (Liu and Inglis, 1992; Le et al., 1995), leading us to postulate that gene 3a might contain an IRES. The predicted structures are five stem-loops (I–V), four of which can be drawn as components of pseudoknots. The free energies of these are calculated to be, respectively, −6.8, +6.0, −3.8, −0.8, and +5.6 kCal/mole by the algorithm of Tinoco et al. (1973), which
A representation of a genetic map with the following features:

- **A**: A diagram showing the T7 transcript with various exons and introns labeled with their respective sizes in kilobases (K).
- **B**: A series of RNA secondary structure diagrams labeled I-V, each showing different nucleotide sequences and interactions.
- **C**: A gel blot analysis with lanes labeled M, ORF7a-3b, ORF7a, Δ11, Δ41, Δ44, Δ120, Δ140, Δ149, Δ289, Δ534, and Δ289-3b. The gel shows bands at 27.7 kDa, 20 kDa, and 7.7 kDa.
- **D**: A table comparing the relative moles of sCAT (7.7, 27.7, and E) for different mutants, including wild type (wt) and deletions (Δ11, Δ41, Δ44, Δ120, Δ140, Δ149, Δ289, Δ534).
are relatively unstable and suggest a low probability for their existence in viral RNA. Nevertheless, to test whether gene 3a might function as an IRES for translation of ORF 3b, deletions within it were prepared and tested. These were (mostly) bidirectional for distances of 16, 86, 74, 192, 314, 337, 334, and 523 nt, and represented 3’-ward deletions of 11, 41, 44, 120, 140, 149, 289, and 534 nt from the first nucleotide in gene 3a, respectively, for which the mutants were named. Results shown in Fig. 6C and summarized in Fig. 6D indicate that the relative molar amounts of sCAT and the 27.7-kDa gene 3b products (~1:0.12) remained essentially unchanged between wild-type and Δ149. The only exception was for Δ41, for which the molar ratio was 1:0.56 along with an inexplicable enhancement of an uncharacterized band with an approximate molecular weight of 6 kDa. For Δ289, which leaves only 17 nt upstream of the ORF 3b start codon, the relative amounts were surprisingly 1:0.07 and not 1:0, as expected. For Δ534, there was abundant synthesis of the sCAT protein but no synthesis of the 27.7-kDa protein; however, there was synthesis of the 20-kDa gene 3b product (O’Connor and Brian, 1999) (Fig. 6D). These results are not consistent with a mechanism of ribosomal entry within gene 3a but rather with one in which ribosomes enter within 50 nt from the start codon of gene 3b. Because the 20-kDa gene 3b product is found with mutant Δ534, the intriguing possibility exists that ribosomes are entering at or downstream of the gene 3b start site and are scanning in the upstream direction to reach the start codon.

**DISCUSSION**

Based on precedents in eukaryotes (reviewed in Mathews, 1996), four mechanistic possibilities should be considered as explanations for how ribosomes approach the downstream ORF 3b on TGEV mRNA 3 for translation: Ribosomes could (1) translate the upstream ORF and then reinitiate synthesis on the downstream ORF, (2) scan through the upstream ORF(s) without the act of translation in a manner known as leaky scanning, (3) bypass the upstream ORF(s) by using an internal ribosomal entry site similar to that used by picornaviruses and flaviviruses on genomic RNA, or (4) bypass the upstream ORF(s) after first binding to the mRNA in a cap-dependent manner and then undergo shunting to a downstream site on the mRNA. Among these, shunting is the most recently recognized and is exemplified by translation on pregenomic RNA of the cauliflower mosaic virus (Futterer et al., 1993) and rice tungro bacilliform virus (Futterer et al., 1996), both pararetroviruses, on adenovirus mRNA (Yueh and Schneider, 1996), and on Sendai paramyxovirus mRNA (Curran and Kolakovsky, 1988, 1989; Latorre et al., 1998).

We conclude that ORF 3b is translated from mRNA 3, and that the likelihood is high that an internal entry of ribosomes is used, possibly one with shuntlike features, and perhaps in conjunction with leaky ribosomal scanning through ORF 3a. The relative contribution of each mechanism on mRNA 3 could not be established by the experiments performed here. However, an internal entry of ribosomes was demonstrated by the use of constructs, in which four extensive ORFs within an 884-nt sequence were placed upstream of ORF 3a and synthesis from ORF 3b was shown to remain approximately one-eighth to one-fifth of that from the 5’-terminal ORF. The internal entry showed some properties of shunting in that (1) no IRES element of the type directing internal entry in picornaviruses and togaviruses could be demonstrated within sequence upstream of gene 3b and (2) translation of ORF 3b in capped transcripts from the synthetic multicistronic psCATORF3a-3b-4 showed some inhibition by a competing soluble cap in the translation mix. That is, internal entry in the multicistronic transcript may follow a cap-dependent step as described for shunting in the adenoviruses, pararetroviruses, and paramyxoviruses.

In general, the mechanistic features of ribosomal shunting, so far described for only viral mRNAs (Jackson, 1996; Mathews, 1996), remain to be clarified. In the case of adenovirus and pararetrovirus mRNAs, an upstream donor structure appears necessary for the shunting step. In pararetrovirus, this appears to be a stable hairpin preceded by a short open reading frame (Hemmings-Miesczak and Hohn, 1999). In the case of TGEV ORF 3b shunting reported here, a requirement for an upstream structure seems unlikely, since shunting took place in the presence of foreign sequence (psCATORF3a-3b-4) as well as (relatively) native sequence (pORF3a-3b-4) at the 5’ terminus. In this respect, the TGEV ORF 3b shunting pathway bears similarity to that in paramyxovirus mRNA, for which no apparent requirement for a donor structure was found (Latorre et al., 1998). Likewise, it is not clear what determines the landing site in a ribosomal shunt. Certainly in the experiments reported here it is not apparent how ribosomes might have been directed to land so close to the 3b initiation codon in TGEV mRNA 3. It is
clearly not the postulated secondary structures within gene 3a, because internal entry took place after these had been removed or disrupted (Δ149, Figs. 6C and 6D). One possibility is that ribosomes are directed to land at or near the start site of gene 3b by specific sequences or by higher-order structures situated very near the landing site. Precedents for this are found in Sendai virus, wherein sequences both upstream and downstream of the Y1 ORF are required for shunt landing (Lattore et al., 1998), and in Hepatitis C virus, wherein sequences extending 28 nt into the ORF are required for IRES-directed landing (Reynolds et al., 1995). Curiously, such a landing site might require that ribosomes backscan to find the gene 3b start codon, a process postulated to explain the translation of certain SV40 and influenza virus transcripts (Peabody et al., 1986; Williams and Lamb, 1989).

Our findings were particularly intriguing because some evidence had suggested the existence of IRES-directed, cap-independent translation for the third ORF in tricistronic IBV mRNA 3 (Liu and Inglis, 1992; Le et al., 1995) and for the second ORF in the bicistronic MHV mRNA 5 (Thiel and Siddell, 1994). The influence of the cap, however, was not examined in the MHV studies, and the possibility remains that a form of shunting might also be exhibited during the translation of these mRNAs.

The consequences of an internal ribosomal entry onto gene 3b for virus replication are not immediately apparent, but one might be that it enables a constitutive synthesis of 3b protein because, in principle, any of the viral mRNAs containing gene 3b (mRNAs 1, 2, and 3) could serve as templates.

MATERIALS AND METHODS

Cells and virus. The Purdue-116 and Miller strains of TGEV were obtained from E. Bohl, Ohio State University. Purdue-116 virus was plaque-purified from infectious genomic RNA, grown on swine testicle (ST) cells in medium containing 10% fetal calf serum (Atlanta Biologicals), and used within eight passages of plaque purification (Sethna et al., 1991; Hofmann et al., 1993). Briefly, cDNA was made from the 5’ end of mRNA 3 with a primer specific to gene 3b (oligo 4(9)) and primers 3968 to 5449 from the genome 3’ end [Fig. 1A; Tung et al., 1992], prepared as described in Kapke et al., 1988,a) into the Psfl site of pGEM-3Z. Second, pSP6ORF3a-3b-4 was created by ligating the 925-bp TGEV Purdue sequences and 4 and 308 nt of gene 5 in vector pGEM-4Z (Promega Biotech) (a sequence obtained from cdNA clone pFT44 [Fig. 1A]), pORF4 (Fig. 1C) was made from pORF3b-4 by first removing the 291-nt M-containing Spfi fragment, religating, and then removing the 716-nt HindIII-BbsI fragment and religating after blunt-ending with mung bean nuclease. pORF3a-3b-4 (Fig. 1C), which carries genes 3a, 3b, and 4 downstream of the T7 RNA polymerase promoter in pGEM-3Z (Promega Biotech), was made in three steps. First, pSP6ORF3a, from which gene 3a and 591 nt of gene 3b can be transcribed with RNA polymerase SP6, was created by ligating the 1111 bp NsiI-PstI fragment from clone pFT39 (a clone containing nucleotides 3968 to 5449 from the genome 3’ end [Fig. 1A; Tung et al., 1992]), prepared as described in Kapke et al., 1988,b) into the Psfl site of pGEM-3Z. Second, pSP6ORF3a-3b-4 was created by ligating the 925-bp BsrGI-EcoRI fragment from pORF3b-4 into the 4048-nt vector-containing EcoRI-BsrGI-linearized fragment of pSP6ORF3a. Third, the entire 1340-nt Spfi-Hinfl fragment inserted from pSP6ORF3a-3b-4, after blunt-ending with T4 DNA polymerase, was ligated in the reverse orientation into similarly blunt-ended EcoRI-SacI-linearized pGEM-3Z.

To place the viral leader upstream of ORF 3a, pLORF3a-3b-4 (Fig. 1C) was constructed by a previously published procedure (Sethna et al., 1991; Hofmann et al., 1993). Briefly, cDNA was made from the 5’ end of mRNA 3 with a primer specific to gene 3b (oligo 4(9)) and primers 3968 to 5449 from the genome 3’ end [Fig. 1A; Tung et al., 1992], prepared as described in Kapke et al., 1988,a) into the Psfl site of pGEM-3Z. Second, pSP6ORF3a-3b-4 was created by ligating the 925-bp BsrGI-EcoRI fragment from pORF3b-4 into the 4048-nt vector-containing EcoRI-BsrGI-linearized fragment of pSP6ORF3a. Third, the entire 1340-nt Spfi-Hinfl fragment inserted from pSP6ORF3a-3b-4, after blunt-ending with T4 DNA polymerase, was ligated in the reverse orientation into similarly blunt-ended EcoRI-SacI-linearized pGEM-3Z.
the overlap procedure to amplify a 1066-nt product from pORF3a-3b-4 DNA. After digestion with restriction enzymes *Nar* and *Bsr* GI, the 932-nt product was cloned into *Nar*- *Bsr* GI-linearized pORF3a-3b-4.

To create pCATORF3a-3b-4 (Fig. 1C), the ORF3a-3b-4-containing 1534-nt *Sph* I fragment from pORF3a-3b-4 was placed into the *Sph* I site of pCAT (Fig. 1C), which was made by cloning the *Bam* HI fragment from pCM4 (Phar-macia) into the *Bam* HI site of *pGEM*-3Z. For better size resolution of the large proteins, the CAT gene in pCATORF3a-3b-4 was truncated by 91 nt on its 3' end by digestions with *Nco* I and nuclease Bal 31, thus forming psCATORF3a-3b-4 (Fig. 1C). This, along with a frameshift, resulted in a total shortening of the CAT protein (now called sCAT) by 46 aa.

The junctions of all constructs were confirmed by sequencing plasmid DNA.

**Preparation of nested deletions within gene 3a, the postulated internal ribosomal entry region.** To obtain deletions within gene 3a, psCATORF3a-3b-4 DNA was linearized at base 7 of ORF 3a with *Thh*111 I, treated with Bal 31 and mung bean nucleases, purified by electrophoresis, and religated. Transformants were screened for deletions by PCR and the sequenced constructs were named for the number of bases deleted downstream of the gene 3a start site (the total number of deleted bases is also noted).

**In vitro translation and analysis of products.** In vitro transcription with T7 RNA polymerase was carried out on linearized plasmid DNAs as recommended (Promega Biotech). pORF3b-4 was linearized with *Eco* RI, pORF3a-3b-4 with *Sca* I, as indicated, and psCATORF3a-3b-4 with Asel. The 1274-nt DNA fragment from *Sca* -linearized pORF3a-3b-4 was purified by affinity chromatography (Geneclean; Bio 101) to ensure transcription of only ORF 3a. For preparation of capped RNA transcripts, 0.5 mM m7G(5')ppp(5')G and 0.25 mM GTP replaced 2.5 mM GTP in the transcription mix (Promega Biotech). Each preparation of RNA was purified by Biospin column chromatography (Bio-Rad), quantitated by spectrophotometry, and monitored for degradation by agarose gel electrophoresis. In *vitro* translation was carried out in methionine-depleted wheat germ extracts or rabbit reticulocyte lysates as recommended by the manufacturers (Promega Biotech and Ambion, Inc.). In some preparations, translation products were treated with RNase A before electrophoresis as recommended by Ambion, Inc., to remove a 27-kDa endogenous band caused by the binding of charged tRNA to proteins. Fifty-microliter reaction volumes contained 50 µCi [35S]-methionine (1000 Ci/mmol; ICN) and 1.0 µg of RNA transcript. To test for inhibition of translation by exogenous methylated cap analog, m7G(5')ppp(5')G (New England Biolabs) was added to the translation mix to final concentrations of 0.1, 0.2, 0.4, and 0.8 mM. Radioactivity in the separated products was quantitated by scanning dried gels with the AMBIS Photoanalytic Imaging System (San Diego, CA) or by scanning autoradiograms of the gels with the Bio-Rad Imaging Spectrophotometer (Bio-Rad). Each experiment depicted was done minimally on three separate preparations of transcript RNA. Standard deviation measurements were made from the results of three separate experiments.

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