trans Regulation of Cap-Independent Translation by a Viral Subgenomic RNA

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Recent years have brought an explosion in discoveries of RNAs that regulate gene expression (3, 33, 38, 56). Many different types of noncoding RNAs posttranscriptionally regulate gene expression in trans (16). These include microRNAs, small interfering RNAs, small nucleolar RNAs, small nuclear RNAs, and bacterial small RNAs (5, 8, 27, 33). trans-acting regulatory RNAs are also generated by viruses. The noncoding adeno virus-associated (VA) RNAs (34) and Epstein-Barr virus-encoded RNAs (EBER) (7) interfere with host antiviral systems and permit efficient expression of late viral genes. Epstein-Barr virus and other herpesviruses also generate microRNAs to downregulate the expression of host and viral genes (38, 44). Red clover necrotic mosaic virus genomic RNA2 directs, in trans, the synthesis of subgenomic mRNA (sgRNA) from genomic RNA1 (52). Flock house virus sgRNA acts as a riboregulator that switches off translation of replication genes from gRNA while permitting translation of structural genes from sgRNA1. These results reveal (i) a new level of control of subgenomic-RNA gene expression, (ii) a new role for a viral subgenomic RNA, and (iii) a new mechanism for RNA-mediated regulation of translation.

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Many positive-strand RNA viruses generate 3’-coterminal subgenomic mRNAs to allow translation of 5’-distal open reading frames. It is unclear how viral genomic and subgenomic mRNAs compete with each other for the cellular translation machinery. Translation of the uncapped Barley yellow dwarf virus genomic RNA (gRNA) and subgenomic RNA1 (sgRNA1) is driven by the powerful cap-independent translation element (BTE) in their 3’ untranslated regions (UTRs). The BTE forms a kissing stem-loop interaction with the 5’ UTR to mediate translation initiation at the 5’ end. Here, using reporter mRNAs that mimic gRNA and sgRNA1, we show that the abundant sgRNA2 inhibits translation of gRNA, but not sgRNA1, in vitro and in vivo. This trans inhibition requires the functional BTE in the 5’ UTR of sgRNA2, but no translation of sgRNA2 itself is detectable. The efficiency of translation of the viral mRNAs in the presence of sgRNA2 is determined by proximity to the mRNA 5’ end of the stem-loop that kisses the 3’ BTE. Thus, the gRNA and sgRNA1 have “tuned” their expression efficiencies via the site in the 5’ UTR to which the 3’ BTE base pairs. We conclude that sgRNA2 is a riboregulator that switches off translation of replication genes from gRNA while permitting translation of structural genes from sgRNA1. These results reveal (i) a new level of control of subgenomic-RNA gene expression, (ii) a new role for a viral subgenomic RNA, and (iii) a new mechanism for RNA-mediated regulation of translation.

BYDV, a major pathogen of wheat and other cereal crops (32), has a positive-sense RNA genome of 5,677 nucleotides (nt) that encodes six ORFs (Fig. 1) (37). Three 3’-coterminal sgRNAs are generated in infected cells. They are not encapsidated and thus are absent at the initial stage of infection. sgRNA1 is the mRNA for the coat protein (CP) (ORF 3), a putative movement protein (ORF 4), and a C-terminal extension of the coat protein required for aphid transmission (ORF 5). Highly abundant sgRNA2 harbors a small ORF (ORF 6) that encodes a predicted polypeptide of 4.3 to 7.2 kDa, depending on the isolate. The sequence of ORF 6 is poorly conserved among BYDV isolates (9), and it is absent in other members of the genus Luteovirus (13, 49). The product of ORF 6 has not been detected in infected cells, although sgRNA2 can be translated in vitro (35). sgRNA3 comprises the 3’-terminal
encodes no ORF, and has no known function. 332 nt of the BYDV genome (26). It accumulates sporadically, cap-independent translation of gRNA and sgRNA1 is conferred by a 100-nt and no poly(A) tail (2). Highly efficient cap-independent trans-

![FIG. 1. trans regulation model of BYDV gene expression. (1) In the early stage of BYDV infection, subgenomic RNAs are absent; thus, the products of ORFs 1 and 2, including the RdRp, are the only proteins produced. (2) Viral genomic-RNA replication and subgenomic-RNA transcription occur. (1, 2, and 3) Viral RNAs accumulate, and viral proteins are produced. (4) The accumulation of sgRNA2 trans inhibits translation of BYDV RdRp from gRNA. (5) However, translation of structural and movement proteins from sgRNA1 is not inhibited. Genomic RNAs are available for encapsidation in the coat protein.

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BYDV genomic RNA (gRNA) and sgRNAs have no 5′ cap and no poly(A) tail (2). Highly efficient cap-independent translation of gRNA and sgRNA1 is conferred by a 100-nt BYDV cap-independent translation element (BTE) in the 5′ untranslated region (UTR) (22, 54). To recruit ribosomes or factors to the 5′ ends of the viral RNAs, where translation initiates, the 3′ BTE must base pair with the 5′ UTR (21). The bridging of the 3′ and 5′ ends of the viral RNAs is facilitated by a kissing stem-loop interaction between the 3′ BTE and a 5-base BTE-complementary loop (BCL) sequence present in the 5′ UTRs of both gRNA and sgRNA1 (21, 48).

In addition to conferring cap-independent translation in cis, the BTE inhibits translation of viral genes and nonviral reporter genes in trans (54, 55). In wheat germ extract, sgRNA2, which harbors the BTE at its 5′ end, trans inhibits translation of gRNA more than that of sgRNA1 (55). The inhibition does not require translation of ORF 6, but it requires a functional BTE in sgRNA2 (55). Premature addition of sgRNA2 at the initial moment of infection strongly inhibits BYDV RNA accumulation (50). Based on these data, the following trans regulation model of gene expression was proposed (Fig. 1). Early in BYDV infection, only the replicase genes, ORF 1 and ORF 2 (translated as a fusion with ORF 1 by ribosomal frameshifting) (4), are translated via BTE-mediated cap-independent translation in cis once replicase is produced, viral RNA is replicated, and gRNA and sgRNAs accumulate. The highly abundant sgRNA2 would selectively inhibit translation of gRNA relative to sgRNA1 in trans. Structural and movement proteins would then be preferentially translated from sgRNA1, and replicase expression would be shut off. Thus, the BYDV viral life cycle would switch from an early to a late stage of gene expression (Fig. 1) (55). Here, we provide evidence that strongly supports the notion that these events occur in virus-infected cells. Moreover, we show that the different translation efficiencies of the BYDV mRNAs in the presence of sgRNA2 are determined by the proximity to the 5′ end of the mRNA of the stem-loop structure that base pairs to the 3′ translation element to allow cap-independent translation. These observations reveal a novel translational control mechanism by a trans-regulatory RNA and a new function for a viral subgenomic RNA.

MATERIALS AND METHODS

Plasmids and RNA constructs. The full-length infectious clone of a PAV isolate of BYDV (BYDV-PAV), pPAV6, was used for transcribing infectious BYDV genomic RNA (11). The sgRNA2 knockout mutant clone of BYDV-PAV, pPAV6SG2, was described previously (28) as SG2G/C. It differs from pPAV6 by a G-to-C mutation at position 4810, which prevents sgRNA2 synthesis. pSG2 and pSG2BF allow T7 transcription of sgRNA2 RNA and its mutant sgRNA2BF, respectively (55). sgRNA2BF contains a GAUC duplication at the BamHI site (BF) of sgRNA2 that destroys the in vitro trans inhibition function of sgRNA2 (55).

Clone pGfLUC was described previously (22) as p5′UTR-LUC-TE869-(A)60-GfLUC is the gRNA reporter transcript derived from SmaI-linearized pGfLUC by in vitro T7 transcription. It encodes the firefly luciferase ORF flanked by the UTRs of BYDV. pRenilla-CP393 was cloned by replacing nt 2643 to 4565 of pPAV6 with the Renilla ORF of pRLuc (Promega, Madison, WI). pSG1LUC was cloned by ligating the Bst1107I-BamHI fragment of pRenilla-CP393 into Bst1107I-BamHI-cut pSG1G, which was described previously (29). SG1LUC is the sgRNA1 T7 transcript from SmaId-cut pSG1LUC. It has the same 5′ UTR as sgRNA1, except for the omission of 14 bases at the extreme 3′ end. pSG1LUC was constructed by replacing the 5′ UTR of pGfLUC with the 5′ UTR of pSG1LUC.

PAV6-FLAG and sgRNA2-FLAG were constructed by inserting a FLAG tag (amino acid sequence, DYKDDDDK) at the 3′ end of ORF 6 in pPAV6 and pSG2, respectively. The FLAG tag insert was created by a three-step PCR approach by using primers bearing a FLAG tag and pSG1 as templates, as described previously (28). The final PCR fragment was cloned into Acc61-Smal-digested pPAV6 or NotI-Smal-digested pSG2. sgRNA2-LIII CS RNA was in vitro transcribed from a PCR-generated template corresponding to the sgRNA2 sequence. The PCR fragment was amplified using LUC 569-LIII-CS as a template and a forward primer bearing a T7 promoter sequence and a reverse primer corresponding to the 3′ end of sgRNA2.

A-GfLUC contained a Ca,C mutation in the loop of stem-loop A (SL-A) and a GAC-to-CUG mutation within the loop of stem-loop D (bases 105 to 107) (see Fig. 5A). First, a BYDV 5′-UTR PCR fragment with a GAC-to-CUG mutation within the loop of stem-loop D was generated by three-step PCR and cloned into NotI-BbsHII-digested pGfLUC. The clone obtained was then used as a PCR template to generate the additional Ca,C mutation. pA-GfLUC was then cloned by ligating the NotI-BbsHIII-digested PCR fragment into NotI-BbsHII-cut pGfLUC. pD-SG1LUC contained an Ap,G mutation in the loop of stem-loop A and an AGUUA-to-CUGACAA mutation of the loop of stem-loop D (bases 111 to 115) (see Fig. 5A). pD-SG1LUC was generated by PCR base mutagenesis of SG1G/C DNA using the Gene Tailor Site-Directed Mutagenesis System (Invitrogen). All constructs were verified by automated sequencing at the Nucleic Acid Facility of Iowa State University.

In vitro transcription and translation. All RNAs were synthesized by using T7 MegaScript kits for uncapped RNA or T7 mMESSAGE mMACHINE kits for capped RNA (Ambion, Austin, TX), according to the manufacturer’s instructions. Prior to transcription, all constructs were linearized either with Smal to generate the same 3′ end as the viral RNA or with VspI to include a 60-nt 3′ poly(A) tail. In vitro translation in wheat germ extract (Promega) and a lucifera
case assay were performed as described previously (50). The reporter RNA transcripts (0.2 pmol), with the indicated molar ratio (n-fold excess) of sgRNA2 or sgRNA2BF transcripts to reporter RNA, were added to the wheat germ translation system with a final reaction volume of 25 μl and translated for 1 h at
25°C prior to luciferase reading. All luciferase assays were performed in at least three independent experiments, each of which was conducted in duplicate or triplicate.

**Western immunoblotting.** Western blotting was performed according to the Amersham Pharmacia Biotech protocol. Total protein from inoculated protoplasts was separated on a 10% polyacrylamide gel and transferred to a polyvinylidene difluoride membrane. After being blocked overnight with phosphate-buffered saline–TWEEN buffer (PTB) containing 5% dried low-fat milk, the membrane was incubated with the primary anti-FLAG antibodies (Sigma, St. Louis, MO) in a 1:25,000 dilution in PTB for 2 h, with a 1:600 dilution of alkaline phosphatase-conjugated anti-fluorescein antibodies for another hour. The membrane was washed three times with PTB after each incubation. After the final incubation with attophase substrate with a volume of 24 μl/cm² for less than 20 min, the membrane was air dried and scanned on a STORM 840 chemiluminescence imager (Molecular Dynamics).

**In vivo translation.** Oat protoplasts were prepared and electroporated with RNA as described previously (12). For the two-step electroporation method, the voltage was reduced to 280 V. For the in vivo translation assay of the SG2fLUC reporter construct, we included a capped, polyadenylated Renilla luciferase reporter RNA as an internal control to normalize electroporation variation. The Renilla luciferase ORF was flanked by the 5’ and 3’ UTRs of the firefly luciferase gene from pGEMLUC (Promega). Luciferase activity was measured 4 h after electroporation.

In the two-step electroporation, oat protoplasts were inoculated with 1 pmol of infectious BYDV PAV6 or PAV6ΔSG2 RNA and incubated for 24 h at room temperature prior to the second electroporation to allow viral replication and sgRNA accumulation. When transfected directly with the nonreplicative sgRNA2 or sgRNA2BF RNA transcripts, the cells were incubated for 4 h prior to the second electroporation. In the second step, protoplasts were inoculated again with 1 to 2 pmol of GfLUC (A-GfLUC), SG1rLUC, or both, as indicated. In all cases, firefly luciferase and Renilla luciferase were analyzed 4 h after the second electroporation, as described previously (51), and the Promega (Madison, WI) Stop-N-Glo system was used to assay both luciferase activities.

**Northern blot analysis.** Total RNAs were extracted from oat protoplasts 24 h postinoculation (p.i.) by using TriZol reagent (Invitrogen, Carlsbad, CA) according to the manufacturer’s instructions. The RNAs were then analyzed by Northern blotting as described previously (29). A 32P-labeled probe complementary to the 1.5-kb 3’ end of BYDV-PAV genome RNA was used to detect BYDV gRNA and sgRNAs (29).

## RESULTS

**Like the actual BYDV RNAs, reporter constructs representing gRNA and sgRNA1 are differentially inhibited by sgRNA2 in trans.** To test the selective inhibition of translation hypothesis diagrammed in Fig. 1, we designed reporter constructs that allowed efficient detection of translation from both gRNA and sgRNA1 in vivo. We replaced the viral coding regions with different luciferase genes that could be assayed in the same tube (Promega Dual Luciferase reporter assay). The reporter construct representing gRNA, GfLUC, includes the firefly luciferase ORF flanked by the BYDV genomic 5’ and 3’ UTRs (Fig. 2A). The reporter construct representing sgRNA1, SG1rLUC, contains the Renilla luciferase ORF in place of ORFs 3 and 4 and most of ORF 5. The 5’ UTR of SG1rLUC contains BYDV nt 2670 to 2842, giving it the same 5’ terminus as sgRNA1. To determine the validity of these two reporter constructs to represent gRNA and sgRNA1, we tested whether GfLUC and SG1rLUC RNAs behave the same as gRNA and sgRNA1 in the presence of sgRNA2 in wheat germ translation experiments (35). When added to translation reaction mixtures containing either GfLUC or SG1rLUC mRNA, sgRNA2 inhibited GfLUC translation more than it inhibited SG1rLUC translation, and this difference increased as the molar ratios of sgRNA2/GfLUC or sgRNA2/SG1rLUC increased (Fig. 2B). The negative control, sgRNA2BF, did not inhibit translation of either reporter RNA (Fig. 2B). sgRNA2BF differs from sgRNA2 only by a GAUC duplication in the BamHI_A site in the BTE. This mutation abolishes both cap-independent translation in cis and inhibition of translation in trans (2, 54).

To determine whether the firefly and Renilla luciferase ORFs caused differences in translation efficiency in the presence of sgRNA2, we constructed an sgRNA1 reporter, SG1fLUC, that differs from GfLUC only in the 5’ UTR (Fig. 2A). SG1fLUC and SG1rLUC behaved indistinguishably in the presence of sgRNA2 (Fig. 2C). Thus, neither the coding regions nor the different lengths of the 3’ UTRs account for the differential effects of sgRNA2 on translation of GfLUC versus SG1rLUC.

To more closely mimic natural infection, gRNA reporter GfLUC, sgRNA1 reporter SG1rLUC, and sgRNA2 were mixed simultaneously in the same wheat germ translation reaction mixture at different ratios. Again, sgRNA2 inhibited translation of GfLUC much more than that of SG1rLUC (Fig. 2D). The difference in translation inhibition by sgRNA2 of GfLUC and SG1rLUC reporters was greater when GfLUC, SG1rLUC, and sgRNA2 were added together in the same translation reaction (compare Fig. 2B and D). The GfLUC/SG1rLUC (gRNA/sgRNA1) expression ratio decreased as the sgRNA2 concentration was increased (Fig. 2E). These results resemble previous observations when gRNA, sgRNA1, and sgRNA2 were all added to a wheat germ translation extract at ratios approximating those in infected cells. In that experiment, gRNA was inhibited by 99% while sgRNA1 was inhibited by only 65%, allowing it to remain a relatively efficient message (55). Thus, the reporter RNAs provide valid representations of the translation of the actual viral RNAs in the presence of sgRNA2. Importantly, these results also reveal that no ORFs or gene products of gRNA or sgRNA1 are necessary for the differential inhibition of translation.

**Differential inhibition of GfLUC and SG1rLUC translation in virus-infected cells.** Having validated that reporter constructs GfLUC and SG1rLUC translate like gRNA and sgRNA1 in the presence or absence of sgRNA2 in vitro, we tested the trans regulation model in oat protoplasts by measuring the translation efficiencies of the two reporter constructs in the presence of replicating BYDV RNA. We employed a two-step electroporation method (Fig. 3A) (50). First, oat protoplasts were transfected by electroporation with the infectious transcript of the BYDV genome, PAV6, or with transcript PAV6ΔSG2. PAV6ΔSG2 RNA has a G4810C point mutation that prevents sgRNA2 synthesis but still permits genomic RNA replication (28). After a 24-hour incubation to allow viral replication and sgRNA accumulation (Fig. 3B), the protoplasts were electroporated again, this time with GfLUC or SG1rLUC reporter RNA. Then after another 4-hour incubation to allow translation of these RNAs, firefly luciferase and Renilla luciferase activities were measured (Fig. 3C and D).

Wild-type and mutant BYDV genomes accumulated to similar levels in protoplasts, with the conspicuous absence of sgRNA2 in protoplasts infected with PAV6ΔSG2 (Fig. 3B). After the second electroporation to introduce reporter mRNA, we observed that the presence of replicating PAV6 RNA caused nearly an 80% drop in the translation of the gRNA reporter GfLUC compared to uninfected cells but caused only a 20% reduction in the translation of the sgRNA1
reporter SG1rLUC (Fig. 3C, PAV6). In cells transfected with PAV6ΔSG2 RNA, translations of GfLUC and SG1rLUC (Fig. 3C, PAV6ΔSG2) were reduced by only 40% and less than 5%, respectively. Thus, infection with PAV6ΔSG2 (which makes no
sgRNA2) inhibited the translation of GfLUC less than did wild-type PAV6 RNA.

When GfLUC and SG1rLUC were coelectroporated in the second step in cells previously transfected with PAV6 RNA, GfLUC translation dropped by 88%, while translation of SG1rLUC actually increased slightly, compared to that of uninfected cells (Fig. 3D, PAV6). SG1rLUC translated at least 10 times more efficiently than GfLUC relative to uninfected cells (Fig. 3D, PAV6). SG1rLUC translation dropped by 70% compared to that of uninfected cells. Thus, the differential inhibition effects of 20-fold molar excess of sgRNA2-LIII-CS, sgRNA2, and sgRNA2BF on translation of GfLUC RNA in wheat germ extract. The error bars indicate standard deviations.

FIG. 4. Differential trans inhibition by sgRNA2 alone in oat protoplasts. The two-step electroporation assay was employed, with 4 h between electroporations. The graph shows the luciferase activities of GfLUC and SG1rLUC measured 4 h after they were coelectroporated with sgRNA2 or sgRNA2BF. The activities of GfLUC and SG1rLUC in cells that were preelectroporated with no RNA were defined as 100%. The error bars indicate standard deviations.

requires a functional BTE but does not require viral replication or infection in vivo.

The mechanism of trans inhibition does not rely on base pairing of the BTE to the 5' UTR of gRNA. A possible mechanism of trans inhibition is base pairing of loop III of the BTE in sgRNA2 with the BCL in the 5' UTR of the gRNA and effective competition with the kissing between the BTE and BCL that occurs in cis to facilitate cap-independent translation. To test this hypothesis, we observed the trans inhibition of the translation of the gRNA in wheat germ extract by a mutant sgRNA2 containing a different sequence in loop III of the BTE that bears no complementarity to the 5' UTR of gRNA. BTE loop III in sgRNA2 was changed from UGUCA to GGCAU (sgRNA2-LIII-CS) and is not complementary to the UGACA in the BCL or any other sequence within the gRNA. sgRNA2-LIII-CS trans inhibited the translation of GfLUC as efficiently as wild-type sgRNA2 (Fig. 5A). In a reciprocal experiment, sgRNA2 with the BF mutation (Fig. 2) did not inhibit in trans, even though it retained complementarity to the gRNA 5' UTR. Thus, the trans inhibition effect of sgRNA2 does not rely on base pairing to the mRNA that it inhibits.

The 5' UTRs of gRNA and sgRNA1 determine the differential trans inhibition effects. The previous results led us to wonder what property of the viral RNA determines differential inhibition. Because neither the ORFs nor the 3' UTRs affect inhibition by sgRNA2 (Fig. 2C), we conclude that features of the different 5' UTRs determine the ability to be inhibited in trans. A striking difference between the two 5' UTRs is that the BCL is in the 5'-proximal stem-loop in sgRNA1 but in the fourth stem-loop (SL-D) from the 5' end in the gRNA (Fig. 6A). The loop of SL-D is located 104 to 109 nt from the 5' end on cap-independent translation in the presence of sgRNA2, we engineered GfLUC so that the 5'-proximal stem-loop of its 5' UTR (SL-A) was able to kiss (base pair to) the BTE, and the natural kissing bases of SL-D were mutated so that it could no longer interact with the 3' BTE (Fig. 6B). This construct, A-GfLUC, translates at an efficiency similar to that of GfLUC (48). To determine whether this relocation of the BCL to the 5'-proximal stem-loop of the gRNA influences the selectivity of trans inhibition by sgRNA2, we tested the translation efficiency of A-GfLUC relative to GfLUC and SG1fLUC in the presence of sgRNA2 in wheat germ extract (Fig. 6C). A-GfLUC behaved similarly...
FIG. 6. Features of the 5' UTRs of gRNA and sgRNA1 that determine the differential *trans* inhibition by sgRNA2. (A) The known secondary structure of the BYDV gRNA 5' UTR (21) and the MFOLD-predicted (58) secondary structure of the sgRNA1 5' UTR. The kissing BCL bases that participate in the long-distance interaction with the 3' BTE are in gray. (B) Schematic diagram of the 3' BTE-5' UTR interactions in the indicated reporter constructs. In A-GfLUC, the 5'-proximal loop of SL-A was converted by a single C-to-A change at position 15 (italics), which made SL-A complementary to the 3' BTE at five consecutive bases (boldface gray). The endogenous SL-D kissing bases were mutated to prevent base pairing with the 3' BTE (GAC to CUG; black italics). In D-SG1fLUC, the loop of sg1SL-D was converted from AGUUA to CUGACAA (bases 110 to 116). The modified D-SG1fLUC also contained an A-to-C change at position 10 in the loop of sg1SL-A, which prevented base pairing to the 3' BTE. (C) Differential effects of sgRNA2 on translation of A-GfLUC, GfLUC, and SG1fLUC in wheat germ extract. The activity of GfLUC, A-GfLUC, or SG1fLUC in the absence of sgRNA2 was defined as 100%. The error bars indicate standard deviations. (D) Differential inhibition by 40-fold excess of sgRNA2 or sgRNA2BF of GfLUC, A-GfLUC, and SG1fLUC translation in oat protoplasts. The two-step electroporation assay was employed as in Fig. 4. GfLUC and SG1fLUC (2 pmol each) were coelectroporated in oat protoplasts 4 h after the indicated RNAs were electroporated into the same cells. SG1fLUC levels differed little in the presence of GfLUC or A-GfLUC, so average SG1fLUC readings are shown. For each reporter RNA, luciferase readings were normalized to the amount detected in the absence of RNA in the first electroporation. (E) Differential effects of 20-fold molar excess of sgRNA2 or sgRNA2BF on translation of GfLUC, SG1fLUC, and D-SG1fLUC in wheat germ extract. The readings were normalized to the amount detected in the absence of sgRNA2 or sgRNA2BF RNA.
to SG1LUC in the presence of various ratios of excess sgRNA2 and thus was inhibited less by sgRNA2 than was GfLUC (Fig. 6C).

To examine in vivo the effect of relocating the kissing loop, a two-step electroporation assay was performed (Fig. 6D). The wild-type (GfLUC) or the modified (A-GfLUC) genomic reporter transcripts were coelectroporated with SG1fLUC RNA in cells previously transfected with sgRNA2 or sgRNA2BF, and the translation efficiencies of both reporter RNAs were compared. While GfLUC translation was inhibited by more than 70% in the presence of sgRNA2, translation of A-GfLUC and SG1fLUC dropped by only 25% and 10%, respectively, compared to mock-transfected cells (Fig. 6D). Thus, unlike the in vitro result (Fig. 6C), A-GfLUC was slightly more susceptible than SG1fLUC to sgRNA2 trans inhibition effects in vivo. Most importantly, A-GfLUC was inhibited far less than GfLUC, from which it differs by only four base changes in the 5’ UTR that allow the BTE to base pair to stem-loop A and not stem-loop D. As expected, sgRNA2BF had little or no inhibitory effect on the translation of any of the reporter RNAs in vitro or in vivo (Fig. 6).

To further analyze the effect of the position of the BCL on selective trans inhibition by sgRNA2, we moved the BCL of SG1fLUC 100 nt downstream to the fourth stem-loop (sg1SL-D) (Fig. 6A) from the 5’ end (construct D-SG1fLUC) (Fig. 6B). We tested the translation efficiency of D-SG1fLUC relative to SG1fLUC and GfLUC in the presence of 20-fold-excess sgRNA2 and sgRNA2BF in wheat germ extract (Fig. 6E). While the 5’-distal position of the BCL in D-SG1fLUC did not affect cap-independent translation in the absence of sgRNA2, D-SG1fLUC was inhibited to a level similar to that of GfLUC in the presence of sgRNA2. As expected, sgRNA2BF did not inhibit the translation of either reporter RNA (Fig. 6E). Taken together, the effects of moving the BCL in either gRNA or sgRNA1 reporters demonstrate that the proximity of the kissing loop to the 5’ end of the RNA is the major determinant of susceptibility to inhibition of translation by sgRNA2.

Lack of a role for sgRNA2 as an mRNA. We next determined whether sgRNA2 functions as an mRNA in cells. sgRNA2 encodes a small, poorly conserved ORF (ORF 6) that is translatable in vitro (55). Unlike gRNA and sgRNA1, sgRNA2 harbors the BTE in its 5’ UTR. No significant sequence complementarity between the 3’ and 5’ UTRs is predicted in sgRNA2. Thus, sgRNA2 is unlikely to be circularized by 5’-3’ base pairing. Because circularization of eukaryotic mRNAs is generally required for efficient translation in vivo but not in vitro (35, 45), we speculated that sgRNA2 may not be translatable in vivo (in protoplasts).

To test the translatability of sgRNA2 in vivo, we fused a FLAG tag to ORF 6 and attempted to detect the P6-FLAG fusion using anti-FLAG antibodies (Fig. 7A). In immunoblots of total protein from cells inoculated with infectious BYDV PAV6, PAV6-FLAG, or nonreplicative sgRNA2-FLAG RNA 24 h posttransfection. As a positive control, the wheat germ translation product of sgRNA2-FLAG is shown in the absence (−) or presence (+) of protoplast extract, which retarded protein mobility in the gel. (C) Northern blot hybridization of total RNA from virus-infected cells showing replication of infectious BYDV PAV6 and PAV6-FLAG. (D) Translation in oat protoplasts of SG2LUC transcript with ORF 6 fused to the firefly luciferase ORF. The presence (+) or absence (−) of a cap and/or a poly(A) tail on this transcript is indicated. Relative luciferase activity was normalized to that of a capped, polyadenylated, nonviral Renilla reporter construct.
A second approach to determine the translatability of sgRNA2 in vivo was to fuse ORF 6 to a luciferase ORF and measure luciferase expression in protoplasts. The 1,800-nt ORF 6-LUC fusion rendered the full-length transcript PAV6 noninfectious, so we examined translation of a transcript representing sgRNA2 containing the ORF 6-LUC fusion (SG21LUC) in cells directly. No luciferase activity was detected representing sgRNA2 containing the ORF 6-fLUC fusion. ORF 6-LUC fusion rendered the full-length transcript PAV6, which we used to measure luciferase expression in protoplasts. The 1,800-nt sgRNA2 in vivo was to fuse ORF 6 to a luciferase ORF and examine the ability of sgRNA2 to fuse ORF 6 to a luciferase ORF, and both reporter RNAs respond the same as viral gRNA and sgRNA1, respectively, in the presence or absence of sgRNA2 in vitro (Fig. 2) (35). This leads to the noteworthy conclusion that neither BYDV coding sequences nor the protein products of the coding regions are necessary for the selective translation of gRNA but has little or no effect on the translation of sgRNA1 in vivo (Fig. 3 and 4). trans inhibition depends on the functional BTE in sgRNA2, as illustrated by the inability of sgRNA2Bf, which differs from sgRNA2 by only a 4-base duplication in an essential region of the BTE, to trans inhibit translation (Fig. 2 to 6). In contrast, trans inhibition does not involve base pairing between the BTE of sgRNA2 and the 5’ UTR of the BYDV reporter RNAs (Fig. 5).

Interestingly, the difference in inhibition by sgRNA2 of translation of gRNA and sgRNA1 reporters was greater in vivo than in vitro. Moreover, these differences were augmented when GfLUC, SG1rLUC, and sgRNA2 were all present simultaneously in vitro and in vivo (compare Fig. 2B and C, Fig. 3C and D). This mixture most closely mimics natural infection and reveals a level of gene regulation in which BYDV RNAs are well coordinated.

A new mechanism of subgenomic-mRNA gene expression control. The data strongly support the trans inhibition model of Fig. 1. Early in BYDV infection, when sgRNAs are absent, only ORF 1 and ORF 2 (replicase genes) are translated via BTE-mediated, cap-independent translation to express the RNA-dependent RNA polymerase. This in turn replicates viral genomic RNA and generates sgRNA1 and particularly large amounts of sgRNA2 (Fig. 3B). The abundant sgRNA2 trans inhibits translation of gRNA only. Unlike in wheat germ extract, where sgRNA1 was inhibited somewhat but to a lesser extent than gRNA, sgRNA1 translation was inhibited only very slightly or not at all in vivo. Thus, the expected large quantity of coat protein can be translated from sgRNA1 as the viral life cycle enters a later stage. This model predicts that the presence of abundant sgRNA2 at the moment of inoculation will block virus replication by prematurely preventing translation of the polymerase, and indeed that was observed previously (50).

We now conclude that there are at least two levels of temporal control of viral-gene expression via subgenomic mRNAs. First and foremost is synthesis of the subgenomic RNAs. They are absent initially; thus, only ORFs 1 and 2 can be translated early in infection from gRNA. sgRNA1 synthesis positively inhibits translation of ORFs 3, 4, and 5. The trans inhibition of translation by sgRNA2 provides the second level of control. It acts negatively to turn off translation of gRNA, favoring translation of sgRNA1 only. Not only does this control the level of RNA-dependent RNA polymerase (RdRp) produced, we propose that this selective trans inhibition could also free the gRNA of ribosomes, making it available for replication and encapsidation.

The proximity of the kissing stem-loop in the 5’ UTR to the 5’ end determines sensitivity to translation inhibition by sgRNA2. The abilities of gRNA and sgRNA1 to be differentially inhibited by sgRNA2 are attributable to their different 5’ UTRs (Fig. 2C). Thus, neither the reporter gene nor the distance of the BTE from the 5’ UTR affected the efficiency of translation in the presence of sgRNA2. In contrast, the mutations that alter the location of the BCL relative to the 5’ end did change the response of the reporter RNA to sgRNA2 (Fig. 6).

The favored translation of sgRNA1 over gRNA in the presence of sgRNA2 may be explained by different structures of the 5’ UTRs per se or by differential requirement for a host factor(s). The positive correlation of proximity of the 5’ UTR kissing stem-loop to the 5’ end and competitiveness of the mRNA in the presence of sgRNA2 (Fig. 6) supports a simple scanning efficiency mechanism. Given evidence that BTE-mediated translation requires 5’-end-dependent ribosome scanning (21), a longer tract, with significant secondary structure, between the 5’ terminus and the BCL may require more translation factors to facilitate ribosome scanning (48). In the absence of sgRNA2, the factors may be in sufficient supply to allow efficient translation of gRNA. Only in the presence of competing sgRNA2 would the factors be reduced enough to hinder translation of gRNA, whereas RNA with the 5’-proximal BCL (sgRNA1) would have a lower factor dependence and hence be less inhibited by competing sgRNA2 (48).

Selective translational control by viral 5’ UTRs in cis has been observed in other viruses. For example, subgenomic RNA 4 of brome mosaic virus has a translational competitive advantage over the other three viral RNAs (46). The 5’ UTR of the coat protein-encoding sgRNA of turnip crinkle virus mediates translation more efficiently than the gRNA 5’ UTR.
likely for and bacterial small RNAs (8, 33, 38). This mechanism is un-
Examples include the microRNAs, small interfering RNAs, of the regulatory RNA to the target RNA(s) to block transla-
by at least two mechanisms. One mechanism is by base pairing gRNA by sgRNA2.
Moreover, it stimulates translation in BYDV 5'-UTR is between two uncapped mRNAs dependent on a 3'-capindependent translation element and is mediated by a third viral RNA (sgRNA2).
Potential mechanism(s) of trans inhibition of translation of gRNA by sgRNA2. Regulatory RNAs inhibit gene expression by at least two mechanisms. One mechanism is by base pairing of the regulatory RNA to the target RNA(s) to block translation or to recruit an inhibitory protein(s) to the target RNA(s). Examples include the microRNAs, small interfering RNAs, and bacterial small RNAs (8, 33, 38). This mechanism is unlikely for trans inhibition by BYDV sgRNA2 because the sequence within the BTE of sgRNA2 that is complementary to the 5'-UTR of the gRNA is not necessary for its trans inhibition activity (Fig. 5). Moreover, it stimulates translation in cis, and also it trans inhibits translation of nonviral mRNAs to which it has no sequence homology (54, 55).
The more likely mechanism is that BYDV sgRNA2 is a molecular decoy that competes for translation initiation factors. In support of this, addition of eukaryotic translation initiation factor 4F (eIF4F) restored translation of mRNA in extracts inhibited by the addition of BTE RNA (54). Indeed, recently eIF4F has been found to interact directly with the BTE and not with the nonfunctional mutant with the filled BamHI site (E. P. Kneller, K. Treder, E. Allen, and W. A. Miller, unpublished data). Thus, eIF4F binding correlates with the trans inhibition function. It is highly unlikely that translation of sgRNA2 is necessary for its function. Not only is no translation product detectable (Fig. 7), but mutant sgRNA2 containing a frameshift mutation that disrupts ORF 6 still selectively inhibited translation of gRNA in vitro (55).
Other trans-regulatory RNAs from viruses have very different functions. Red clover necrotic mosaic virus RNA2 has a 34-nt trans activator sequence, which is required for transcription of sgRNA from RNA1 (52) and for encapsidation (6). This trans activator base pairs to RNA1 to facilitate sgRNA synthesis. Flock house virus sgRNA3, but not its translation product, trans activates replication of viral genomic RNA2 (14, 15), which then down-regulates the synthesis of sgRNA3 from genomic RNA1 (57). Epstein-Barr virus EBER RNAs may function similarly to VA RNAs in blocking the host protein kinase RNA-activated antiviral response, because they can re-
cue replication of adenovirus lacking VA RNAs (7, 10). The herpesvirus microRNAs downregulate both viral and host gene expression at different stages of the viral infection (38, 44). Among the above-mentioned viral regulatory RNAs, VA RNAs, EBER RNAs, and most likely BYDV sgRNA2 are not mRNAs. Others function as both a coding RNA and a noncoding regulatory RNA.
It remains to be investigated whether other viruses employ this type of selective, negative regulation of translation. Competitive inhibition of translation among RNAs of other multi-
trans inhibition of translation of one viral RNA but not another, (ii) trans inhibition of cap-independent translation, and (iii) inhibition by an apparently untranslated RNA all are properties that so far are known only in BYDV infection. It will be particularly interesting to know whether other viruses that generate multiple subgenomic mRNAs, such as severe acute respiratory syndrome coronavirus and other members of the Nidovirales, or the Closteroviridae, are also controlled by a subgenomic RNA in trans. By extension, it is possible that a host mRNA with particularly high affinity for translation factors could negatively regulate translation of other host mRNAs by this mechanism.

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