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Gennadiy Koev
Iowa State University

B. R. Mohan
Iowa State University

W. Allen Miller
Iowa State University, wamiller@iastate.edu

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Primary and Secondary Structural Elements Required for Synthesis of Barley Yellow Dwarf Virus Subgenomic RNA1†

GENNADIY KOEV, B. R. MOHAN,‡ AND W. ALLEN MILLER*

Plant Pathology Department, Iowa State University, Ames, Iowa 50011-1020

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Barley yellow dwarf luteovirus (BYDV) generates three 3′-coterminal subgenomic RNAs (sgRNAs) in infected cells. The promoter of sgRNA1 is a putative hot spot for RNA recombination in luteovirus evolution. The sgRNA1 transcription start site was mapped previously to either nucleotide 2670 or nucleotide 2769 of BYDV genomic RNA (gRNA) in two independent studies. Our data support the former initiation site. The boundaries of the sgRNA1 promoter map between nucleotides 2595 and 2692 on genomic RNA. Computer prediction, phylogenetic comparison, and structural probing revealed two stem-loops (SL1 and SL2) in the sgRNA1 promoter region on the negative strand. Promoter function was analyzed by inoculating protoplasts with a full-length infectious clone of the BYDV genome containing mutations in the sgRNA1 promoter. Because the promoter is located in an essential coding region of the replicate gene, we duplicated it in a nonessential part of the genome from which a new sgRNA was expressed. Mutational analysis revealed that secondary structure, but not the nucleotide sequence, was important at the base of SL1. Regions with both RNA primary and secondary structural features that contributed to transcription initiation were found at the top of SL1. Primary sequence, but not the secondary structure, was required in SL2, which includes the initiation site. Disruption of base pairing near the sgRNA1 start site increased the level of transcription three- to fourfold. We propose that both primary and secondary structures of the sgRNA1 promoter of BYDV play unique roles in sgRNA1 promoter recognition and transcription initiation.

Many positive-strand RNA viruses express genes via RNA-template transcription of subgenomic mRNAs. Several mechanisms of subgenomic RNA (sgRNA) synthesis have been proposed for various viruses, including internal initiation of the replicate at subgenomic promoters, premature termination of negative-sense RNA synthesis with subsequent independent replication, 5′ leader-primed synthesis, and RNA recombination (reviewed in references 19 and 27). So far, only internal initiation on the negative-strand template has been demonstrated as a mechanism of sgRNA synthesis in plant RNA viruses (16, 29, 47), although premature termination during negative-strand synthesis has been suggested as an alternative mechanism (27, 43). Despite the lack of direct evidence for internal initiation of transcription in most viruses, we will adhere to convention and refer to cis elements responsible for synthesis of sgRNAs as sgRNA promoters.

Boundaries of sgRNA promoters have been determined for several RNA viruses in vivo (3, 4, 15, 17, 23, 45, 47, 48). Their sizes vary from 24 nucleotides (nt) in Sindbis virus (23) and 27 nt in cucumber necrosis virus (17) to nearly 100 nt in beet necrotic yellow vein virus (3). With the exception of beet necrotic yellow vein virus (3), the larger portions of subgenomic promoters are located upstream of the transcription initiation site (in the positive sense).

In vitro analysis of the sgRNA4 promoter of brome mosaic virus (BMV) and its comparison with other alpha-like virus subgenomic promoters revealed four major structural elements: a core promoter, an AU tract downstream of the initiation site, an oligo(A) tract, and an enhancer element located upstream of the core (24). However, wild-type levels of RNA4 synthesis in vivo require an additional upstream element that contains repeats of sequences from the core (15). The core promoter of BMV RNA4 has been characterized extensively in vitro, revealing sequence requirements for transcription initiation and suggesting that the primary and not the secondary structure of RNA is critical for specific and accurate initiation, much like in RNA-dependent RNA polymerase promoters (42). However, RNA secondary and tertiary structures play important roles in various processes of the virus life cycle including RNA replication, recombination, translation, and others. These processes involve RNA-protein interactions (1, 6, 10, 35, 38). The combination of primary and secondary RNA structure requirements is common in control of virus replication (21, 36, 38, 40). Therefore, we set out to determine the role of RNA structure in sgRNA promoter function.

The object of this study is the PAV strain of barley yellow dwarf virus (BYDV), which belongs to the genus Luteovirus (formerly called subgroup I) of the family Luteoviridae (formerly the luteovirus group) (9, 26). BYDV has a positive-sense, 5.7-kb genomic RNA (gRNA). In the infected cell, three 3′-coterminal sgRNAs are produced (12, 18). They are not encapsidated. All three sgRNAs play different roles in virus replication. sgRNA1 is the mRNA for the coat protein, a readthrough extension of the coat protein involved in aphid transmission, and a 17-kDa protein required for plant systemic infection (8). sgRNA2 codes for a small 4.3- to 7.0-kDa peptide that is dispensable for virus replication in protoplasts (33). sgRNA2 may also regulate translation from gRNA and sgRNA1. The role of sgRNA3 which lacks open reading frames (ORFs) is unclear (27, 28, 31).

The family Luteoviridae is split into two major genera, Luteovirus and Polerovirus, based on differences in the 5′ halves of
TABLE 1. Primers used to construct mutants of the duplicated sgRNA promoters

<table>
<thead>
<tr>
<th>Construct</th>
<th>5′ primer (sequence)</th>
<th>3′ primer (sequence)</th>
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<tr>
<td>pPAV6G1A</td>
<td>(ATAGGTACCCGAGGTGGTTGAGTCGTCAC)</td>
<td>SL (ATAGGTACCAATGTTGAGTCGTCACC)</td>
</tr>
<tr>
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<td>SL (ATAGGTACCAATGTTGAGTCGTCAC)</td>
<td>SL (ATAGGTACCAATGTTGAGTCGTCAC)</td>
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<td>SL (ATAGGTACCAATGTTGAGTCGTCAC)</td>
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<tr>
<td>pJ</td>
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<td>SL11A (ATAGGTACCAATGTTGAGTCGTCAC)</td>
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* KpnI recognition sequences are underlined, mutant nucleotides are in boldface, and deletions are shown as underlined spaces.

their genomes (9, 26). The genes in these regions, including the RNA-dependent RNA polymerase (RdRp), are unrelated between the genera (28). The border between divergent and homologous regions is located between ORF2 and ORF3 (RdRp and coat protein genes) (28). This region also includes the 5′ end of sgRNA1. Based on this observation, we proposed that recombination has occurred during luteovirus evolution by replicase strand switching at the subgenomic promoters (28, 30).

As a first step in testing the recombination model, we have begun mapping the sgRNA promoters of BYDV. In this study, we mapped the primary and secondary structures required for sgRNA synthesis in detail. We show that both primary and secondary RNA structure play unique roles in promoter recognition by viral replicase in vivo.

MATERIALS AND METHODS

Plasmids. pPAV6 is a full-length cDNA clone of BYDV-PAV described in reference 11. pGK1 was constructed by cloning the Avr1 (2456)-Ssp1 (2737) fragment of pPAV6 into pGEM-3Z digested with Avr1 and Ssp1. Mutants pKel, pKel-f, and pKel-M were constructed by two-step PCR (20). To make pKel-f and pKel-sh in the first round of PCR, we used an upstream mutagenic primer, 5′-GGCATTCAATGTTGAGTCGTCAC-3′ (altered bases in boldface), spanning bases 2655 to 2690 and a downstream primer 5′-TGGGCACTTACCGTAACACTATAAGTACCACCTTCA-3′ (altered bases in boldface), spanning bases 2663 to 2684 and the above downstream primer. The product of the first round was gel purified and used in the second round of PCR as the downstream primer with the upstream primer

TACGACCCCACTACGACGGTCTTAC)

TTACGACCCCACTACGACGGTCTTAC)

TACGACCCCACTACGACGGTCTTAC)

TTACGACCCCACTACGACGGTCTTAC)

TTACGACCCCACTACGACGGTCTTAC)

TTACGACCCCACTACGACGGTCTTAC)

TTACGACCCCACTACGACGGTCTTAC)

TTACGACCCCACTACGACGGTCTTAC)

TTACGACCCCACTACGACGGTCTTAC)

TTACGACCCCACTACGACGGTCTTAC)

TTACGACCCCACTACGACGGTCTTAC)

TTACGACCCCACTACGACGGTCTTAC)

TTACGACCCCACTACGACGGTCTTAC)
Australia and Illinois isolates were used by Kelly et al. and by Dinesh-Kumar et al., respectively. However, the high homology of the two isolates as well as a conserved hexanucleotide, GUGAAG, present at the 5′ ends of the gRNA and sgRNA1 and sgRNA2, revealed in the study by Kelly et al. (18), prompted us to revisit this issue. We constructed a probe, pGK-1, that is complementary to the region of BYDV gRNA (bases 2456 to 2737) that spans the sgRNA1 start site as mapped by Kelly et al. (2670) but should not detect sgRNA1 as mapped by Dinesh-Kumar et al. (2769 [Fig. 1]). This probe hybridized with sgRNA1 of BYDV-PAV-Illinois and an in vitro transcript that contains the 5′ end at 2670. It did not detect an in vitro transcript with the 5′ end at 2769. Therefore, the 5′ extremity of sgRNA1 of this isolate is in fact well upstream of 2769 and consistent with the 5′ end at 2670 in BYDV-PAV-Australia (Fig. 1). Promoter mapping described herein supports an initiation site at or near 2670.

**Mutations near the sgRNA1 start site affect transcription.** The sgRNA1 initiation site (2670) is located within ORF2, which encodes viral RdRp (Fig. 1), which is required for replication (33). To examine the possibility of using deletion mutagenesis to map the subgenomic promoter, we introduced a stop codon at position 2650 of the infectious clone, pPAV6, which truncated ORF2 by 30 3′-terminal codons. No replication of this mutant transcript in oat protoplasts was detected by Northern blot analysis (data not shown). Thus, deletion mapping of the subgenomic promoter in this region was not possible.

To determine the importance of individual nucleotides around the start site for sgRNA synthesis, we introduced point mutations in this region of pPAV6. Two mutants with five base changes around the start site replicated but synthesized no sgRNA1 (Fig. 2). One had a Val→Asp amino acid change; the other had no amino acid changes (Fig. 2). To test the importance of the G at the initiation site, it alone was mutated to a C in mutant 2670M. This resulted in an unavoidable amino acid substitution (Val→Leu). This mutant did not replicate at all (Fig. 2). This is surprising considering that the more radical change of Val→Asp at this site did not knock out replication. Thus, either the RdRp tolerated Asp but not Leu at amino acid position 825, or the particular base that coincides with the 5′ end of sgRNA1 is essential for gRNA replication. Mutants Kel-6 and Kel-f demonstrated the sensitivity of sgRNA1 transcription to changes in the conserved hexanucleotide GUG

**RESULTS**

**Reexamining the 5′ end of sgRNA1.** The 5′ end of sgRNA1 of BYDV-PAV was mapped to position 2769 of gRNA by Dinesh-Kumar et al. (12) and to position 2670 by Kelly et al. (18). The 99-nt discrepancy has been attributed to the two different isolates of BYDV-PAV used in these studies: Australia and Illinois isolates were used by Kelly et al. and by
AAG and immediately upstream of the initiation site. Mutant 2670M further emphasized the difficulty of the promoter characterization due to potential undesired alterations in RdRp function.

Mapping the boundaries of the sgRNA1 promoter. To allow more mutagenesis of the sgRNA1 promoter, we moved a copy of it to a nonessential portion of the genome. This duplication of subgenomic promoters and this synthesis of sgRNAs from ectopic locations have been demonstrated elsewhere for several RNA viruses (3, 4, 15, 22, 45, 47). We duplicated a 314-nt region flanking the putative sgRNA1 initiation site (nt 2503 to 2816) and introduced it into a unique Kpn1 site in ORF5 (position 4154 [Fig. 3A]). This ORF is not required for virus replication in protoplasts (33). As expected, the promoter duplication resulted in the expression of an additional 1.7-kb sgRNA (sgRNA1A [Fig. 3B]). This caused a dramatic drop in accumulation of sgRNA1 from its natural setting and also reduced gRNA levels. This phenomenon of reduced synthesis of sgRNAs from upstream promoters when additional promoters are inserted downstream has been observed elsewhere with other RNA viruses (15, 47). Reduced gRNA accumulation is likely due to lack of coat protein synthesis (33) caused by reduced levels of its mRNA (sgRNA1). This ectopic expression of sgRNA1 set the stage for the deletion mapping and detailed characterization of the sgRNA1 promoter without interfering with the RdRp coding region.

We tested a set of mutants containing various portions of the 314-nt promoter region for sgRNA1A synthesis in oat protoplasts. The smallest construct capable of directing sgRNA1A transcription, 2SL, consisted of the 98-nt RNA sequence from bases 2595 to 2692. Three smaller constructs, I (nt 2611 to 2692), J (nt 2595 to 2679), and K (nt 2611 to 2679), were incapable of producing the artificial sgRNA1A (Fig. 3C). The 2SL construct did not cause such a large reduction in sgRNA1 and gRNA accumulation as did construct PAVSG1A, perhaps because it was not as strong a promoter as the 314-nt insert. Furthermore, the 314-nt duplication in PAVSG1A may have given a gRNA too large to be encapsidated, while the smaller overall size of 2SL gRNA (98-nt duplication) may have permitted encapsidation.

Secondary structure prediction for the sgRNA1 promoter. To explore the possible role of RNA secondary structure in sgRNA1 synthesis, we analyzed the 98-nt promoter sequence for potential RNA folding patterns by using the MFOLD program (49). Because sgRNA synthesis has been shown to occur by internal initiation of transcription on the negative-strand template in other viruses, we used the complement of the mapped subgenomic promoter region for the secondary structure predictions. Most of the suboptimal folding patterns contained two stem-loop structures (SL1 and SL2). There were two major variations of the SL1 folding: structure I and structure II (Fig. 4A). To establish which one is more likely to exist, we compared the sgRNA1 promoter regions of other BYDV isolates and the related soybean dwarf virus (SbDV). The BYDV sequences were too highly conserved to shed light on the secondary structure, while SbDV diverged significantly (Fig. 4B). The predicted structure of the SbDV subgenomic promoter region resembled only that of BYDV-PAV structure II. Sequence covariations found in SbDV revealed four sites at which base changes retained base pairing (boxed base pairs [Fig. 4A]) in structure II. Thus, structure II is more likely to exist in BYDV, even though it is calculated to be slightly less stable than structure I (Fig. 4A).

Comparison of the sgRNA1 promoter with those of sgRNA2 and sgRNA3 and the 3' end of the negative strand of gRNA did not show any significant sequence homology except for the
earlier identified hexanucleotide 3'-CACUUC-5' in sgRNA1 and sgRNA2 and gRNA (5'-GUGAAG-3' in the positive sense). RNA secondary structure predictions of the negative-strand RNA in those regions revealed a hairpin with the initiation site in its stem similar to SL2 (Fig. 4C). No structure like SL1 was predicted in the other sgRNA promoter regions. This suggested that SL2-like structure may be a common structural element of RdRp recognition regions.

**Nuclease probing of the sgRNA1 promoter secondary structure.** To test the existence of the two computer-predicted stem-loops in the sgRNA1 promoter, we constructed a plasmid, pT7SGP1, which contained a region of the negative-strand gRNA spanning the minimal sgRNA1 promoter (nt 2595 to 2716) cloned behind the bacteriophage T7 promoter. For RNA secondary structure analysis, we used 5'-end-labeled in vitro transcripts of the sgRNA1 promoter obtained from KpnI-linearized pT7SGP1. To detect double- and single-stranded RNA regions within the subgenomic promoter, we performed partial digests of the 5'-end-labeled transcripts with RNase T1 (cuts single-stranded G's) and imidazole, a chemical RNase that cleaves RNA at all single-stranded bases. Imidazole has been used elsewhere to resolve secondary structure of various RNAs (14, 46).

Both the T1 and the imidazole analyses identified most of the single-stranded RNA regions corresponding to those predicted by computer (Fig. 5A). The lower part of SL2 was sensitive to nuclease, suggesting the existence of a single-stranded junction between the two major stem-loop domains. The upper and the lower regions of SL1 were well protected from both T1 and imidazole digestion, implying stable RNA helices. Highly protected C2648/U2649/A2650 and U2602/G2603/G2604 appeared paired to each other, whereas both G2605 and G2647 were nuclease sensitive and consistent with the bulge in structure II (Fig. 4A and 5A). Therefore, despite the lower predicted stability, structure II appears more likely to form in solution than does structure I.

The middle portion of SL1 exhibited ambiguous base pairing. Both the base-paired and the single-stranded conformations may coexist in dynamic equilibrium (breathing), reflecting the weak base pairing of the AU-rich AUUCU:AGAAU helix. Based on their nuclease sensitivities, both terminal loops of SL1 and SL2 seemed well defined and consistent with the computer prediction (Fig. 5A). The RNase probing analysis superimposed on the phylogenetically conserved, computer-generated secondary structure allowed us to propose the solution structure of the sgRNA1 promoter (Fig. 5B).

**Primary and secondary RNA structures of SL1 are required for transcription of sgRNA1.** To test the involvement of primary and secondary RNA structure elements of SL1 in sgRNA synthesis, we introduced a series of mutations into the duplicated sgRNA1 promoter. Oat protoplasts were inoculated with BYDV.

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**FIG. 4. RNA sequence and secondary structure analysis of the sgRNA1 promoter of BYDV.** (A) Two secondary structures (I and II), with the calculated free energies predicted with MFOLD (49), contain two stem-loops, SL1 and SL2. Boxes indicate covariations that preserve the predicted secondary structure. The right-angled arrow indicates the initiation site (nt 2670). (B) Alignment of RNA sequences in the subgenomic promoter regions of five BYDV strains and SbDV. The BYDV strains are PAV-Australia (also known as PAV-Vic) (pav-aus), PAV-Japan (pav-jap), PAV-Purdue (pav-p), MAV (mav), and MAV (mav). The bottom row shows consensus sequence (cons). Dashes indicate bases that do not differ from consensus. (C) Computer-predicted stem-loop structures in the genomic and subgenomic sgRNA1 promoter regions of BYDV. The sequence is negative sense; numbering is positive sense. The conserved hexanucleotides at the initiation sites are in boldface.
the mutants, and total RNA was isolated 24 h postinoculation (hpi) and analyzed by Northern blot hybridization. We used the ratio of steady-state levels of sgRNA1A to those of gRNA as a measure of the promoter activity.

To examine the role of the helix at the base of SL1, we introduced nucleotide alterations (blocks of 4 nt) in both strands of the helix that disrupted and restored the base pairing (mutants SL11A, SL11B, and SL11C [Fig. 6A]). Transcription of sgRNA1A was eliminated when the base pairing was disrupted in mutants SL11A (sgRNA1A/gRNA ratio = 0.06 ± 0.02) and SL11B (0.02 ± 0.01) and was restored to a level higher than that of wild type (construct 2SL, 0.51 ± 0.08 [Fig. 6B]) in the compensatory mutant SL11C (1.56 ± 0.25 [Fig. 6B]). These results indicated that the secondary structure, and not the nucleotide sequence at the bottom of SL1, is required for transcription.

To test the role of the upper part of SL1, we introduced similar mutations into the upper helix (SL12A, SL12B, and SL12C [Fig. 6A]). Both mutants SL12A and SL12B, which disrupted the base pairing, exhibited low levels of sgRNA1A accumulation (0.12 ± 0.05 and 0.11 ± 0.02, respectively [Fig. 6B]). The compensatory mutant SL12C failed to restore the promoter activity (0.05 ± 0.02 [Fig. 6B]), indicating that specific nucleotide sequences on both sides of the helix (and possibly RNA secondary structure as well) are important in this region.

To examine the role of the single-stranded and ambiguous regions of SL1 in sgRNA1A synthesis, mutant SL1D was constructed with two sequences tracts deleted: U2602-A2620 and C2641-G2657 (Fig. 6A). Surprisingly, this mutant produced low but significant levels of sgRNA1A (0.10 ± 0.01 [Fig. 6B]), indicating that the middle portion of SL1 could be deleted while SL1 still retained a low level of transcription. Deletion of the bulged U2635 (SL1U [Fig. 6A]) reduced transcription (0.19 ± 0.04 [Fig. 6B]), indicating its importance, but not its absolute requirement, for the promoter activity. Changing the sequence of the terminal loop of SL1 to its complement (Fig. 6A) yielded low levels of sgRNA1A (mutant SL13, 0.10 ± 0.03 [Fig. 6B]). Thus, the specific sequence of the terminal loop is also important for transcription.

Nucleotide sequence but not the secondary structure of SL2 is required for transcription of sgRNA1. We next characterized the sequence and structural elements of SL2 that are involved in sgRNA synthesis. As reported for initial mutants, sequence alterations within five bases of the transcription start site in its natural location in ORF2 knocked out sgRNA1 synthesis (Fig. 2). Mutations in Kel-6, Kel-f, and 2670M were predicted to disrupt the secondary structure of SL2. To separate the influence of the nucleotide sequence alteration from

FIG. 5. Nuclease probing of the sgRNA1 promoter secondary structure. (A) Imidazole and T1 RNase partial digests of 5'-end-labeled transcript of pT7SGP1 containing the sgRNA1 promotor region (negative sense). Gel-purified, end-labeled RNA was incubated in 0 M (0) and 0.8 M (I) imidazole under non-denaturing (native) conditions for 17 h (lanes 3 and 4). The nondenaturing T1 digest was performed with 0.01 U of the enzyme for 5 min at 37°C (lane 5) (see Materials and Methods). Denaturing digests with the T1 (cuts after G) and U2 (cuts A → U) RNases generated markers in lanes 1 and 2. The products were separated on a 6% polyacrylamide gel containing 8 M urea. The straight line beside lane 4 indicates predicted base-paired regions; dashed lines represent predicted single-stranded junctions and ambiguous regions; curved lines show predicted loops and bulges. Double and single arrows represent G's that were cleaved strongly and weakly, respectively, by T1 nuclease. Filled circles indicate uncut or very weakly cut G's. (B) Solution structure of the sgRNA1 promoter. Arrowheads represent the T1 analysis data; triangles represent the imidazole digestion data. Larger and smaller symbols indicate strong and weak cuts, respectively.
that of the RNA secondary structure disruption, we designed a
series of sgRNA1 promoter mutants that disrupted and re-
stored the secondary structure of SL2 in the duplicated pro-
moter. Three nucleotides near the start site were altered to
disrupt base pairing in either strand of SL2 in mutants trpl1
and trpl2 (Fig. 7A). The mutant trplc contained both of the
above sets of substitutions to restore the structure of SL2 (Fig.
7A). No sgRNA1A synthesis was detected in trpl1 (0.01 ±
0.02) and trplc (0.0 ± 0.02), both of which are mutated in the
conserved hexanucleotide at the start of sgRNA1 (Fig. 7B).
However, trpl2 exhibited a level of transcription three times higher than that of wild type (1.60 ± 0.02). Thus, the primary structure adjacent to the start is required for sgRNA synthesis (Fig. 7B), and SL2 secondary structure may actually inhibit transcription.

In order to examine the role of the initiating C (G2670 in the positive strand), we changed it to a G (mutant 2670G [Fig. 7A]). We also weakened the bottom of SL2 by changing the complementary G to a C (2688C [Fig. 7A]). In the double mutant, comp1, the base pairing was restored while the alteration of the primary structure was maintained (Fig. 7A). The results were similar to those in the previous experiment: no transcription in 2670G (−0.02 ± 0.01) and comp1 (−0.16 ± 0.21), and transcription almost four times higher than the wild-type level of transcription in 2688C (1.95 ± 0.32) (Fig. 7B).

Finally, changing the sequence of the SL2 terminal loop to that of its complement in the mutant SL21 (Fig. 7A) reduced the sgRNA1A level to 0.11 ± 0.01 (Fig. 7B), indicating the importance, but not the absolute requirement, of the nucleotide sequence of the SL2 terminal loop. These results stressed the importance of the primary RNA sequence and the negative effect of the secondary structure at the initiation site.

In pursuit of the nucleotides in SL2 that could be altered without affecting transcription, we replaced the 5′-terminal conserved hexanucleotide of sgRNA1 (GUGAAG in the positive strand) with the 5′-terminus of sgRNA3 (GACGAC in the positive strand). Interestingly, this mutant, SL2SG3 (Fig. 7A), did not produce detectable levels of sgRNA1A (0.0 ± 0.02 [Fig. 7B]). We also constructed a mutant, SGP300, with the duplicated putative sgRNA3 promoter region (301 nt, 5150 to 5450) spanning the sgRNA3 start site (nt 5348) inserted in the KpnI site in ORF5 in order to test if this promoter could function outside its wild-type location. The mutant produced sgRNA1A (0.72 ± 0.04 [Fig. 7B]), which indicated that the sgRNA3-specific 5′-terminal hexanucleotide could function in the context of its own promoter. This shows that the replicase recognizes very different promoters.

**DISCUSSION**

**Rerevaluation of the 5′ end of sgRNA1.** Here we show that the 5′ end of sgRNA1 is most likely at position 2670 as reported by Kelly et al. (18) and not at our originally reported site of 2769 (12). The error may have occurred due to a variety of technical difficulties, involving mismatches between the probe (from BYDV-PAV Australia) and the viral RNA (BYDV-PAV Illinois), and the unexpectedly far-upstream location of the 5′ end. The difficulty of mapping 5′ ends of *Luteoviridae* sgRNAs is further indicated by discrepancies reported for potato leafroll virus. Initially, the 5′ end of sgRNA1 of potato leafroll virus was reported to be only 40 nt upstream of the coat protein start codon (44). However, subsequent analysis mapped it to 212 nt upstream at a region which, like BYDV (nt 2670 to 2675), shows homology to the 5′ end of the gRNA (25). Phylogenetic comparisons support this latter start site for other members of the *Luteoviridae* (28).

**Structure and function of sgRNA1.** Mutations in the subgenomic promoter unveiled roles for three types of structures: (i) one in which the secondary and not the primary structure is important (base of SL1), (ii) an ambiguous region where the primary structure and possibly secondary structure both may influence the transcription efficiency (upper stem and loop of SL1), and (iii) a region where the primary and not the secondary structure is required (stem of SL2). The nucleosome probing and phylogenetic analysis support the existence of the base of SL1. Stem-loops have been predicted in sgRNA promoters of other RNA viruses (24, 47, 48), with the initiation site usually located within a single-stranded region. To our knowledge, our data represent the first actual demonstration of a requirement for a specific helical domain in a viral sgRNA promoter. The requirement for primary and not secondary structure at the sgRNA1 initiation site result is consistent with other studies showing the role of single-stranded regions for specific protein recognition (2, 35, 36).

The ambiguous results of the structure probing of the distal portion of SL1 lead us to propose that this portion of the subgenomic promoter forms metastable structures. Alternative conformations have been demonstrated for other viral RNAs (32, 39); therefore, a portion of molecules may also fold as predicted in structure I, or the entire AUUCU:AGAAU stem of structure II may be unpaired, giving a very large bulge between stems at the top and the base of SL1. The 36-base deletion mutant (SL1D) lacking this ambiguously paired region still retained 20% of wild-type promoter activity. This indicates that the deleted region probably does not contact the viral replicase directly but may provide favorable spatial localization of the essential elements.

**Recognition of the sgRNA1 promoter by replicase.** We propose that SL1 acts as a replicase recognition site, placing the replicase in proximity to the start site at the base complementarity to 2670. The double-stranded proximal end (bottom) of SL1 may serve only to provide the structural foundation that ensures that the sequence at the distal end (top) of SL1 (Fig. 6) is presented in the proper orientation for specific binding by the replicase. The ambiguously structured, nonessential, central portion of SL1 may play only an auxiliary role in spacing between the distal region and the initiation site.

This model of a separate RNA binding site and adjacent initiation site resembles those proposed for a recombination site in TCV satellite RNAs, subgenomic promoter recognition by BMV replicase (42), and recognition of bacteriophage Qβ RNA by its replicase (5). The TCV satellite RNA sequence includes a bulged stem-loop as the putative replicase binding site adjacent to the actual site at which RNA synthesis takes place (35). The well-characterized BMV promoter differs by its lack of requirement for the secondary structure in replicase recognition (42). However, this difference may be explained in part by the use of a cell-free transcription system which may have less stringent requirements for *cis* elements in vitro than those observed in vivo (15). The divergent sequences of the three BYDV promoters may allow differential expression of each. Each sgRNA promoter may have a separate recognition site on the replicase holoenzyme, each of which may be on a separate protein factor as shown for positive- and negative-strand recognition by Qβ replicase (5), or the sequences may have different affinities for the same site on the replicase.

**Possible alternative mechanisms of sgRNA1 synthesis.** Our data leave open the possibility that sgRNAs could be synthesized by premature termination during negative-strand synthesis on genomic template followed by independent replication of the sgRNA (27). Evidence suggesting such a mechanism has been provided for coronaviruses (7) and dianthoviruses (43). Upstream of the sgRNA1 5′ end, stem-loops complementary to SL1 and SL2 are predicted to exist in the positive strand by MFOLD (49). The mutations that support a role for the helix at the base of SL1 in the negative strand could equally well support the role of the complementary helix in the positive strand. Such a structure in the positive strand could inhibit replicase migration along the template, favoring termination. The resulting 3′ end of the truncated negative strand would resemble that of full-length negative strand owing to the CAC UUC homology, allowing it to be recognized and replicated by...
the replicase. Thus, that essential sequence would still be serving the promoter function.

The premature termination model has been invoked for red clover necrotic mosaic dianthovirus (RCNMV), owing to a remarkable base pairing between the positive strands of the two gRNAs of the virus that is essential for formation of sgRNA from gRNA1 (43). The polymerase of RCNMV is closely related to that of BYDV (29, 31), and so a similar replication mechanism might apply to BYDV. Because BYDV has only one gRNA, the termination structure would form either as the complement of SL1, as discussed above, or intermolecularly, in which two gRNA molecules dimerize by base pairing at the complementary sequences that would otherwise form the stem-loop. Alternatively, it is possible that other inter- or intramolecular interactions could generate a transcription termination structure.

Some observations argue against a premature termination model. The disruption of the secondary structure at the initiation site (SL2) that increases sgRNA1A synthesis (Fig. 7) should have had the opposite effect if sgRNA1A was synthesized by premature termination during the negative-strand synthesis because stable stems should increase termination (37). Furthermore, the role of SL1 is more than just providing a stem-loop structure, which would (in the positive strand) block replicase migration, because mutations at the distal end of SL1 reduced sgRNA1A accumulation independently of their effect on secondary structure.

Role in recombination. The presence of an sgRNA promoter at the 3’ end of ORF2 is consistent with our proposed model in which Luteoviridae genomes recombine at a subgenomic promoter in the vicinity of the intergenic region between ORF2 and ORF3 (28). This model requires that gRNAs are generated by internal initiation of the replicase on the negative strand. A premature termination mechanism, followed by independent replication of the sgRNA, is difficult to reconcile with the sgRNA promoter being a recombination hot spot, if a stem-loop in the positive strand facilitates termination. However, if base pairing between gRNAs occurs as with RCNMV, one could imagine replicate occasionally switching gRNA strands, rather than terminating at this base-paired region. This type of recombination, mediated by base pairing between template strands, has been demonstrated for BMV (34).

Evolution of sgRNA promoters. The small sizes of viral genomes often require the overlapping of protein coding regions with cis-acting RNA elements. It is an intriguing question, how genetic information coding for a protein and an RNA cis element with which it interacts coevolved on the same region of viral genome. ORF2 is very sensitive to deletions and point mutations (Fig. 2), while the sgRNA promoters tolerate changes and consist of quite diverse sequences. Thus, we propose that sgRNA promoters evolved independently at the appropriate genomic locations while allowing overlapping ORFs to maintain their function. The size of the BYDV sgRNA1 promoter is comparable to that in other RNA viruses, but no apparent sequence homology can be found with subgenomic promoters of members of other virus groups. This diversity among sgRNA promoters of related virus taxa and ability to tolerate movement to different regions of the genome (3, 4, 15, 22, 45, 47) further support the hypothesis of multiple, independent origins of sgRNA promoters. The only conserved secondary structures among the BYDV promoters are the SL2-like hairpins flanking all initiation sites (Fig. 4C), yet the SL2 stem structure inhibits sgRNA1 transcription (Fig. 7). Perhaps the SL2-like stems serve as negative regulatory elements to prevent too much transcription of the sgRNAs at inappropriately stages in RNA replication. Obviously, much additional research is necessary to unveil this complex interplay of replicase-RNA interactions.

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