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The authors note, “The rRNA loading panels in Figs. 1D and 2D were inadvertently duplicated by Olivier Voinnet. Part of the image in Fig. 2D was also used in Figs. 3C and 5C, as explained in the respective legends. The experiments for Figs. 2, 3, and 5 were all run on the same agarose gel and blotted onto the same filter, to allow their direct comparison through the common reference dilution series seen in Figs. 2D, 3C, and 5C. The samples in Fig. 1D were run and blotted independently and it is therefore likely that the rRNA loading image in Fig. 1D, but not that in Fig. 2D, is erroneous. We no longer have the original files used for this paper and we are not able to provide the correct rRNA loading control to Fig. 1D. We recognize, therefore, that the equal loading of the samples in the figure is now not supported by the presented data. However, we are confident that the conclusions from the figure about the geminiviral suppressor AC2 are correct, as that interpretation was later confirmed in multiple independent publications.”

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Suppression of gene silencing: A general strategy used by diverse DNA and RNA viruses of plants

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In transgenic and nontransgenic plants, viruses are both initiators and targets of a defense mechanism that is similar to posttranscriptional gene silencing (PTGS). Recently, it was found that potyviruses and cucumoviruses encode pathogenicity determinants that suppress this defense mechanism. Here, we test diverse virus types for the ability to suppress PTGS. *Nicotiana benthamiana* exhibiting PTGS of a green fluorescent protein transgene were infected with a range of unrelated viruses and various potato virus X vectors producing viral pathogenicity factors. Upon infection, suppression of PTGS was assessed in planta through reactivation of green fluorescence and confirmed by molecular analysis. These experiments led to the identification of three suppressors of PTGS and showed that suppression of PTGS is widely used as a counter-defense strategy by DNA and RNA viruses. However, the spatial pattern and degree of suppression varied extensively between viruses. At one extreme, there are viruses that suppress in all tissues of all infected leaves, whereas others are able to suppress only in the veins of new emerging leaves. This variation existed even between closely related members of the potexvirus group. Collectively, these results suggest that virus-encoded suppressors of gene silencing have distinct modes of action, are targeted against distinct components of the host gene-silencing machinery, and that there is dynamic evolution of the host and viral components associated with the gene-silencing mechanism.

In plants, posttranscriptional gene silencing (PTGS) is manifested as the reduction in steady-state levels of specific RNAs after introduction of homologous sequences in the plant genome. This reduction is caused by an increased turnover of target RNA species, with the transcription level of the corresponding genes remaining unaffected (reviewed in ref. 1). Recently, it was shown that PTGS involves systemic spread of a silencing signal directing sequence-specific RNA degradation (2, 3). Although the exact mechanism by which PTGS operates has yet to be elucidated, various findings that viruses can both initiate and be targets of PTGS (4) led to the suggestion that PTGS is a natural mechanism by which plants recognize and combat foreign nucleic acids (5).

In support of the proposed relationship between PTGS and virus resistance, it was shown that some viruses induce an RNA-mediated defense (RMD) in nontransgenic plants. This induced defense is similar to PTGS in that it is characterized by nucleotide sequence-specific resistance against virus infection (6). In some but not all instances, the upper leaves of plants exhibiting this RMD were said to have recovered because they contained only low levels of viral RNA and were symptom-free (7, 8).

However, the ability of viruses to infect plants indicates that they have evolved to avoid or suppress the RMD. This idea was first prompted by analysis of potyviral synergistic interactions with other viruses (9). It was shown that this synergism was the result of suppression of a host defense mechanism by the Hc-protease (HcPro) encoded in the potyviral genome (10). Subsequent studies further established that HcPro was a suppressor of PTGS and provided a link between PTGS and antiviral defense (11–13). Presumably, the suppression acts against the RMD evoked above. A second protein, the 2b protein of cucumber mosaic virus (CMV), was also identified as a suppressor of PTGS in *Nicotiana benthamiana* (12). Interestingly, HcPro and the 2b proteins did not target the silencing mechanism in the same way; HcPro suppressed silencing in tissues where it was already established, whereas the 2b protein only affected silencing initiation (12, 14).

Although 2b and HcPro are dissimilar at the protein sequence level, they are both pathogenicity determinants of their respective viruses (15, 16). By extrapolation, we predicted that many viral pathogenicity determinants would be identified as suppressors of gene silencing and that, more generally, many viruses would have the ability to suppress PTGS. Here, we test these ideas by infecting *N. benthamiana* plants exhibiting PTGS of a green fluorescent protein (GFP) transgene with a range of viruses. Plants were also infected with potato virus X (PVX) vectors expressing previously identified viral pathogenicity determinants. If these wild-type and recombinant viruses produced suppressors of a PTGS-like resistance mechanism, we predicted that they would interfere with PTGS of GFP. The outcome of these experiments was consistent with our prediction and revealed that suppression of gene silencing is a widespread strategy among plant viruses. Our study led to the identification of three viral suppressors of PTGS and revealed an intriguing phenotype of silencing suppression that operates in the vicinity of the veins.

**Materials and Methods**

**Plant Material.** Transgenic *N. benthamiana* plants carrying the GFP ORF were described previously (12).

**PTGS Suppression Assay.** Leaves of seedlings of line 16c were infiltrated with a strain of *Agrobacterium tumefaciens* carrying a binary Ti plasmid vector, into which a functional 35S-GFP cassette had been inserted, as reported (17). After 15–20 days, when PTGS of GFP was achieved in the whole plant, a systemic leaf was inoculated with a wild-type or recombinant virus. This leaf is referred to as “inoculated leaf.” The challenged virus was then allowed to spread in the silenced plant, and two types of leaves were collected at 14 or 20 days postinoculation (DPI). “Old leaves” were infected leaves that had emerged before the virus had spread systemically, whereas “new leaves” were those emerging after the virus had moved systemically.

**Wild-Type Viruses.** Isolates of alfalfa mosaic virus (AMV), foxtail mosaic virus (FoMV), narcissus mosaic virus (NMV), nandina virus X (NVX), viola mosaic virus (VMV), and tomato bushy stunt virus (TBSV) were obtained from Roger Hull from the John Innes Centre (JIC) collection (Norwich, U.K.). Cowpea mosaic virus (CPMV) was obtained from George Lomonosoff at JIC. African

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Abbreviations: GFP, green fluorescent protein; PTGS, posttranscriptional gene silencing; RMD, RNA-mediated defense; HcPro, Hc-protease; DPI, days postinoculation; PVX, potato virus X; CMV, cucumber mosaic virus; PKV, potato virus Y; ACSV, African cassava mosaic virus; TBSV, tomato bushy stunt virus; TMV, tobacco mosaic virus; CPMV, cowpea mosaic virus; RYMV, rice yellow mottle virus; NMV, narcissus mosaic virus; NVX, nandina virus X; VMV, viola mosaic virus; FoMV, foxtail mosaic virus; OL, old leaves; NL, new emerging leaves.

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Table 1. Suppression of PTGS of GFP mRNA caused by various plant viruses

<table>
<thead>
<tr>
<th>Virus group</th>
<th>Virus</th>
<th>Suppression of PTGS</th>
<th>Old leaves/New leaves</th>
<th>Whole leaf/Vein centric</th>
<th>Protein*</th>
<th>Other known functions†</th>
</tr>
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<tbody>
<tr>
<td>Alfamovirus</td>
<td>ALMV</td>
<td>0/9</td>
<td>—</td>
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<td>?</td>
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<tr>
<td>Comovirus</td>
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<td>Geminivirus</td>
<td>ACMV</td>
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<td>OL and NL</td>
<td>Whole leaf</td>
<td>AC2</td>
<td>Virion sense gene expression transactivator</td>
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<tr>
<td>Nepovirus</td>
<td>TBRV</td>
<td>0/6</td>
<td>—</td>
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<td>Potexvirus</td>
<td>PVX</td>
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<td>—</td>
<td>—</td>
<td>?</td>
<td>—</td>
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<tr>
<td>FoMV</td>
<td>0/9</td>
<td>—</td>
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<tr>
<td>NMV</td>
<td>8/9</td>
<td>OL and NL</td>
<td>Whole leaf</td>
<td>?</td>
<td>—</td>
<td>—</td>
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<tr>
<td>NVX</td>
<td>7/9</td>
<td>OL and NL</td>
<td>Whole leaf</td>
<td>?</td>
<td>—</td>
<td>—</td>
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<tr>
<td>MVV</td>
<td>7/9</td>
<td>OL and NL</td>
<td>Whole leaf</td>
<td>?</td>
<td>—</td>
<td>—</td>
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<tr>
<td>Potyvirus</td>
<td>PVY/TEV</td>
<td>10/10</td>
<td>OL and NL</td>
<td>Whole leaf</td>
<td>HcPro</td>
<td>Genome amplification</td>
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<td></td>
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<td>Viral synergism</td>
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<tr>
<td>Sobemovirus</td>
<td>RYMV</td>
<td>—‡</td>
<td>—‡</td>
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<td>P1</td>
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<tr>
<td>Tobamovirus</td>
<td>TMV</td>
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<td>OL and NL</td>
<td>Vein centric</td>
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<td>—</td>
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<tr>
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<td>TRV</td>
<td>7/9</td>
<td>OL and NL</td>
<td>Whole leaf</td>
<td>?</td>
<td>—</td>
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<tr>
<td>Tombusivirus</td>
<td>TBSV</td>
<td>7/9</td>
<td>NL only</td>
<td>Vein centric</td>
<td>19K</td>
<td>Host-specific spread and symptom determinant</td>
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</table>

PTGS of the GFP mRNA was induced in transgenic *N. benthamiana* by Agrobacterium infiltration, as described (17). After systemic infection, suppression of gene silencing was assessed under UV illumination over time and confirmed by RNA gel blot analysis. RNA samples were taken from either old leaves that had emerged before the virus had spread systemically (OL) or new leaves emerging after virus infection (NL). The total number of plants tested is indicated as well as the phenotype of suppression in leaves (affecting whole tissues or vein centric). Viruses were tested in duplicate independent experiments during the summer and the winter.

*The identification of the 2b protein and HcPro as PTGS suppressors is described in refs. 11–13. The identification of AC2, P1, and the 19K protein as PTGS suppressors is described in this study.

†Appropriate references can be found in refs. 16 (2b), 21 (AC2), 13 and 15 (HcPro), 24 (P1), and 22 [19K (19-kDa)].

‡RYMV is not infectious in *N. benthamiana*. The P1 protein has been identified as a PTGS suppressor by expression from the PVX vector.

Recombinant Viruses. The P1 protein sequence of an RYMV isolate from Nigeria (20) was amplified by using the following 5′ phosphorylated primers: ATG ACT CGG TTG GAA GTT C-3′ for the intact protein (P1) and ATC ACA CGG TTG TAA GGTT TC-3′ for an untranslatable protein (mP1). The phosphorylated downstream primer used for amplification was CAT CCC GTG TCA GTG C (19). The two PCR fragments were cloned into the EcoRV site of the PVX vector (p2C2S) (19). The orientation of RYMV PCR fragments was confirmed by colony-PCR using antisense primer in the vector sequence at the 3′ end of the p2C2S multiple cloning site (GTA GTG GAG GTA GTT GCC CC) and the two sense RYMV 5′ primers described above. PVX-AC2 and PVX-mAC2 (21) were provided by John Stanley. PVX-HS142 and PVX-HS160 (22) referred to as PVX-19k and PVX-m19k, respectively, were provided by Andrew Jackson, University of California, Berkeley.

In Vitro Transcription and Northern Blot Analysis. In *in vitro* transcription reactions to produce infectious recombinant PVX RNAs and inoculation were as described (19). Northern blot analysis was described previously (12).

GFP Imaging. Visual detection of GFP was as described (12). Close-up images were obtained by using a Leica MZ FLIII dissecting stereomicroscope coupled to a fluorescence module. The filter set used for GFP imaging was the GFP-plus fluorescence set from Leica (excitation 480 nm, dichromatic beam splitters, 505 nm, Barrier filter 510 nm). Photographs were produced by using a Leica MPS60 device coupled to the stereomicroscope.

Results

Suppression of Gene Silencing by Diverse Plant Viruses. A test for silencing suppression was based on a previously described experimental system (12). This system involves transgenic *N. benthamiana* plants carrying a highly expressed GFP transgene that makes them fluoresce bright green under UV illumination. Systemic silencing in these plants was induced by infiltration of lower leaves of transgenic seedlings with a strain of *A. tumefaciens*, as described (17). By 20 days postinfiltration, silencing of the GFP was extensive in all vegetative tissues of the plants, and, consequently, they appeared uniformly red under UV illumination. At this stage, there was no PTGS in the growing points of the plant, and silencing was maintained by being constantly initiated in nonsilenced cells located near or in the meristems (17). These silenced plants were then infected with a range of plant viruses, and, when systemic symptoms were observed, the extent of green fluorescence was assessed under UV illumination. In addition, Northern blot analysis was performed to assess the level of GFP mRNAs in infected tissues.

Our findings were that many but not all of the viruses tested suppressed gene silencing in *N. benthamiana* (Table 1). With several viruses, suppression occurred in old leaves (OL) that had emerged before the virus had spread, as well as in new emerging leaves (NL). This was reminiscent of the pattern of silencing suppression previously described for PVY (12). In contrast, TBSV suppressed gene...
silencing only in new emerging tissues, as was previously reported for CMV (12, 14). FoMV, alfalfa mosaic virus, or tobacco black ring virus were like PVX in that they were fully infectious but did not have any effect on GFP silencing. From the diversity of viruses tested in this analysis, we conclude that PTGS suppression is a property of many plant viruses. However, because the spatial pattern and degree of suppression varied extensively between viruses, it was likely that different mechanisms would be involved.

The Geminivirus-Encoded AC2 Protein Is a Suppressor of Gene Silencing. As shown in Table 1, infection of ACMV led to suppression of GFP silencing at about 3 wk postinoculation in both fully expanded and new emerging infected tissues (Fig. 1B). Correspondingly, Northern blot analysis revealed that GFP mRNA levels were high in both types of tissues and that suppression also occurred in inoculated leaves, although to a lower extent (Fig. 1D, lanes 1–4). Therefore, these results were consistent with a suppressor of PTGS encoded in the ACMV genome.

To identify this putative suppressor, we exploited previous findings that a PVX vector expressing the AC2 protein (PVX-AC2) produced necrotic symptoms that were much more severe than those of wild-type PVX, suggesting that AC2 suppressed a host defense mechanism (21). From the above results, it was likely that AC2 was a suppressor of RMD.

The test of this hypothesis was to infect GFP-silenced plants with PVX-AC2 (Fig. 1A). As a control, plants were also inoculated with PVX-mAC2 (Fig. 1A) in which a single point mutation introduces a premature stop codon in the AC2 ORF (21). At about 2 wk postinoculation, PVX-AC2-infected plants exhibited severe symptoms, as reported (21). Under UV illumination, most of the infected tissues, including leaves that had emerged prior to virus inoculation, were green fluorescent (Fig. 1C), and GFP mRNA levels were similar to those in nonsilenced GFP plants (Fig. 1D, lanes 5 and 6). In contrast, PVX-mAC2 did not produce severe symptoms and did not suppress GFP silencing (Fig. 1D, lanes 7–8). From these results, we conclude that the AC2 protein encoded in the ACMV genome is a suppressor of maintenance of PTGS in N. benthamiana.

Vein-Specific Suppression of Silencing by the 19-kDa ("19K") Protein of TBSV. N. benthamiana infected with TBSV showed reversion of PTGS at about 3 wk postinoculation, when symptoms were fully systemic (Table 1). As in CMV-infected plants, the restoration of green fluorescence occurred only in new emerging infected leaves. However, this suppression of silencing was weaker than with CMV, so that the green fluorescence was barely detectable under UV illumination from a hand-held lamp. Also unlike CMV, TBSV suppressed PTGS only in and around the veins (Fig. 24). Vein-specific reversion of GFP was more evident when detached, new emerging leaves were observed under a dissecting microscope (Fig. 2D). Northern blot analysis showed that GFP RNAs were more abundant in the new leaves of the infected plants than in old leaves or in mock-inoculated, nonsilenced plants. However, the GFP RNA in the new leaves was <20% of the level in mock-inoculated plants (Fig. 3D, lanes 4 and 5).
Other Examples in Which Suppression of PTGS Occurs Preferentially in or Near the Veins. As part of our survey, we investigated the effect of TMV and CPMV, type members of the tobamovirus and comovirus groups, respectively. Inoculation of the corresponding viruses onto GFP-silenced plants led to suppression of gene silencing that affected both new emerging and already expanded silenced tissues, thus indicating that maintenance of PTGS was alleviated (Table 1, and Fig. 3A and B). However, as shown previously for TBSV and PVX-19K, suppression was mostly manifested near or in the veins with most tissues of the lamina remaining silenced (i.e., red fluorescent), although symptoms of the respective viruses were observed on the whole leaf lamina (data not shown). This phenotype did not change over time, even when infected leaves were fully expanded and completely infected. With both viruses, green fluorescence in the vicinity of the veins was very strong, and this effect was clearly apparent under UV illumination from a hand-held lamp (Fig. 3A and B). Northern blot analysis of RNAs extracted from infected leaves showed that GFP RNA accumulation was restored in those tissues but at a low level when compared with the abundance of GFP RNA extracted from similar tissues of nonsilenced, noninfected plants (Fig. 3C and D). This was probably because of dilution of the vein tissue into the most abundant silenced tissues of the lamina. Therefore, this molecular analysis was consistent with the particular phenotype of silencing suppression observed under UV illumination.

It has been reported that the 19K protein of TBSV is a pathogenicity determinant. For example, a PVX vector expressing the 19K protein (pHS142), referred to here as PVX-19K (Fig. 1A), induced severe symptoms on N. benthamiana (22). In addition, inactivation of the 19K protein in TBSV had an attenuating effect on the lethal apical necrotic symptom phenotype that is usually elicited in plants by TBSV (22). Collectively, these data indicate that the TBSV 19K protein possesses attributes of a suppressor of gene silencing. To test this hypothesis, silenced GFP plants were inoculated with PVX-19K (Fig. 1A). As a control, plants were also inoculated with pHSl60 (referred to here as PVX-m19K) carrying a nontranslatable form of the 19K protein (Fig. 1A, and ref. 22). By 2 wk postinoculation, plants infected with PVX-19K exhibited very severe symptoms, whereas PVX-m19K-infected plants had mild mosaic symptoms, as reported (22). Suppression of silencing occurred in PVX-19K-infected plants, but was manifested only in new emerging tissues and was most pronounced in the veins (Fig. 2B). However, symptoms of PVX-19K were visible on all areas of the leaves (data not shown). Similar tissues infected with PVX-m19K remained uniformly red fluorescent (Fig. 2C). Northern blot analysis of RNA extracted from new emerging, infected leaves showed that only low levels of GFP RNAs could be detected in PVX-19K-infected tissues (Fig. 2D, lanes 6 and 7) and that GFP RNAs were below the level of detection in PVX-m19K-infected tissues (Fig. 2D, lanes 8 and 9). Taken together, these results suggest that the 19K protein of TBSV is a weak suppressor of PTGS in N. benthamiana that operates in the vicinity of the vein tissues of new emerging leaves.

The inocula of these related viruses had been quantified using the local lesion host Chenopodium amaranticolor (25) and diluted, so that they would be comparable to a PVX inoculum used as an internal reference (40 lesions per leaf). Following infection, we confirmed that these viruses gave similar types of symptoms. Thus, the variation in the suppressor of silencing activity reflected intrinsic properties of the viruses rather than the degree of infection. Surprisingly, the variable suppressor activity did not correlate with the nucleotide sequence similarity of these viruses. PVX and
We predicted that many viruses would encode proteins that are suppressors of an RMD mechanism and that these proteins would also suppress PTGS (12). The likely candidate suppressors were viral proteins that, like the 2b protein or HcPro, were originally characterized as pathogenicity determinants. Consistent with this hypothesis, the ACMV AC2, the RYMV P1, and the TBSV 19K pathogenicity factors all suppress PTGS of a GFP transgene. It is therefore likely that the activity of these proteins in pathogenicity of the encoding virus is associated with suppression of RMD. The ability of these proteins to enhance symptoms of PVX vectors is most likely explained in the same way. The finding that a DNA geminivirus, ACMV, encodes a suppressor was not surprising because other geminiviruses are known to induce PTGS, and presumably RMD, in transgenic and nontransgenic plants (26, 27).

Each virus produced a characteristic pattern of silencing suppression. Some, like potyviruses, suppressed in young and old leaves. Others were like CMV and affected only young leaves. There was also variation in the tissue specificity with ACMV, VMV, NMV, NVX, and PVX-P1 affecting all tissues, whereas TBSV, TMV, and CPMV specifically suppressed silencing in tissues that were in or close to the veins. We do not think that these differences reflect the tissue tropism of these viruses. Similar patterns were reproduced when various suppressors were expressed from a PVX vector that has been shown to express foreign proteins uniformly throughout infected leaves (19). A more likely explanation depends jointly on the mode of action of the suppressor and the component of the gene-silencing mechanism that is targeted. For example, if a suppressor can degrade a component required for maintenance of gene silencing, it will have an effect in both new and old leaves. However, if the suppressor blocks synthesis or activation of a component required for silencing, the suppression would be restricted to new emerging leaves, where silencing would be established in the presence of the viral suppressor. In old leaves, the component would have been formed in the absence of the suppressor and, consequently, would be unaffected when the virus would infect the plant.

The suppression of silencing in veins, for example with the 19K protein of TBSV, could indicate that this protein is stable or expressed only in the veins or that it is targeted against a component of the PTGS mechanism that is qualitatively or quantitatively different between vascular and nonvascular tissue. Alternatively, the suppressor could be targeted against the systemic signal of PTGS. We have shown that this signal is phloem-transmitted and that, in recipient leaves, it is primarily located in and near the veins (17). Of these alternative explanations for suppression of silencing in veins, we consider that those involving vein-specific components

![Image of plants infected with PVX-P1](image-url)
or stability of the suppressors are unlikely because, in all cases, PTGS suppression extended into cells outside the vascular bundle and appeared to reflect movement of the signal rather than a precisely vein-specific silencing process. For this reason, we propose that the suppressors of TMV, CPMV, and TBSV are likely targeted against the systemic signal of silencing and may therefore represent a viral adaptation to systemic RMD.

Although TMV, TBSV, and CPMV are able to suppress PTGS only in or near the veins, they are nevertheless able to accumulate at a high level throughout the infected leaf. It is likely, therefore, that these viruses have secondary strategies for counteracting the effects of RMD. These strategies may involve evasion, so that the process is not activated, or escape from the antiviral mechanism. Luteoviruses, which are typically restricted to the phloem (25), may provide an interesting example of viruses that are unable to either suppress, evade, or escape from the effect of RMD outside the veins. Consistent with this idea, it has been reported that the level of potato leafroll luteovirus (PLRV) increased up to 12-fold in *Nicotiana* species that were coinfected with NMV, tobacco rattle virus (TRV), or PVY (28). It now seems likely that this increase was due, at least in part, to the ability of PLRV to spread beyond the veins as a result of suppression of RMD in the double-infected plants. Here, we show that NVM, TRV, and PVY are all able to suppress maintenance of PTGS in *N. benthamiana* (Table 1). In contrast, coinfected with alfalfa mosaic or tobacco black ring viruses that are unable to suppress PTGS (Table 1) did not alter PLRV concentration in leaves (28).

**Gene Silencing Activation/Suppression as a Coevolutive Mechanism?**

It is striking that the viral suppressors of silencing are so diverse. So far, we have been unable to identify any common structural features in these proteins, and we conclude that the suppressor function has evolved independently several times as a strategy to counteract the effects of RMD. In some instances, it is conceivable that some suppressors have converged toward the same function and, thus, are targeted against similar components of the silencing machinery. For example, the RYMV P1 protein shares striking functional similarities with the potyviral-encoded HcPro protein. First, when produced from the PVX vector, both proteins are suppressors of maintenance of PTGS in *N. benthamiana*. In addition, both proteins are required for efficient accumulation of viral RNA in protoplasts and long distance movement in their respective host (15, 24).

Because RMD is likely to have a central role in plant–virus interactions, one can also anticipate that there will be a dynamic evolution of plant components required for the mechanism and, accordingly, of the virus-encoded components necessary to overcome it. The poty- and potexvirus groups probably represent different stages in this dynamic evolution. In the potyvirus group, the HcPro of tobacco etch virus (TEV) (11, 13), PVY (12), and pea seedborne mosaic virus (O.V., unpublished data) are suppressors of GFP silencing in *Nicotiana* species. In these viruses, the suppressor seems to be a conserved function, and its corresponding target is also likely to be conserved in different plants. Similarly, the target of the RYMV P1 protein may be conserved from rice to tobacco. In contrast, the potexvirus strategy for counteracting RMD and PTGS is apparently in a state of evolutionary flux. Presumably, PVX and FoMV, as opposed to VMV and NVX, do not have a functional suppressor of silencing in *N. benthamiana* and, on that host, must use alternative strategies to escape or evade the mechanism, as proposed above. However, it might be expected that on other hosts, PVX and FoMV would produce functional suppressors and, conversely, VMV and NVX would not. The test of this coevolution hypothesis would require a suitable set of host plant species exhibiting PTGS, rather than the single GFP *N. benthamiana* line used here.

In due course, it may transpire that the balance between RMD activation and suppression will strongly influence virus–host interactions (29). For example, if a virus cannot suppress, evade, or escape the effects of the mechanism, the inoculated plant will be considered as a nonhost because there will only be subliminal infection. Similarly, if the virus is able to suppress the mechanism but cannot block the signal of silencing, it is likely that local or systemic spread of the virus will be impaired. Probably the best prospect for understanding this proposed adaptive process involves characterization of mutants impaired in PTGS (30) and identification of host components interacting with viral suppressors. In addition, the increasing body of evidence that PTGS also operates in animals raises the fascinating possibility that silencing suppression has also been adopted by animal viruses (31).

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