Heterologous RNA-silencing suppressors from both plant- and animal-infecting viruses support plum pox virus infection

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INTRODUCTION

RNA silencing is a general term that refers to a complex set of RNA-guided gene-regulatory mechanisms controlling crucial physiological processes, such as developmental patterning, responses to stress conditions and maintenance of genome stability, in a wide variety of eukaryotic organisms (Ambros & Chen, 2007; Baulcombe, 2005). Among its functions, RNA silencing plays a major role in natural antiviral immunity at least in plants, fungi and invertebrate animals, where infecting viruses induce the production of small interfering (si)RNAs from viral dsRNAs and/or secondary RNA structures by the action of RNase-III ribonuclease Dicer-like (DCL) proteins. These virus-derived (v)siRNAs are then incorporated into RNA-induced silencing complexes (RISCs), guiding them, by base-pair complementarity, to viral RNAs for degradation.

HCPo, the RNA-silencing suppressor (RSS) of viruses belonging to the genus Potyvirus in the family Potyviridae, is a multifunctional protein presumably involved in all essential steps of the viral infection cycle. Recent studies have shown that plum pox potyvirus (PPV) HCPo can be replaced successfully by cucumber vein yellowing ipomovirus P1b, a sequence-unrelated RSS from a virus of the same family. In order to gain insight into the requirement of a particular RSS to establish a successful potyviral infection, we tested the ability of different heterologous RSSs from both plant- and animal-infecting viruses to substitute for HCPo. Making use of engineered PPV chimeras, we show that PPV HCPo can be replaced functionally by some, but not all, unrelated RSSs, including the NS1 protein of the mammal-infecting influenza A virus. Interestingly, the capacity of a particular RSS to replace HCPo does not correlate strictly with its RNA silencing-suppression strength. Altogether, our results suggest that not all suppression strategies are equally suitable for efficient escape of PPV from the RNA-silencing machinery. The approach followed here, based on using PPV chimeras in which an under-consideration RSS substitutes for HCPo, could further help to study the function of diverse RSSs in a ‘highly sensitive’ RNA-silencing context, such as that taking place in plant cells during the process of a viral infection.

Intriguingly, it still remains controversial whether this mechanism is part of the innate antiviral response in vertebrate animals (for a complete review of RNA silencing-based viral immunity systems, see Ding, 2010).

In turn, to counteract this defensive response, viruses have evolved a variety of strategies, the most common of which is the expression of proteins that block the RNA-silencing machinery of the host. These factors are called RNA-silencing suppressors (RSSs) and show a large diversity in amino acid sequence and anti-silencing mechanisms (Burgyán & Havelda, 2011; Dunoyer & Voinnet, 2005; Roth et al., 2004; Valli et al., 2009). The helper component–protease (HCPo) of plant potyviruses was the first RSS to be described (Anandalakshmi et al., 1998; Brigneti et al., 1998; Kasschau & Carrington, 1998). It is a potent suppressor that blocks RNA silencing by hijacking specifically 21 nt long vsiRNAs (Lakatos et al., 2006) and, possibly, by other mechanisms (Endres et al., 2010). HCPo is a cysteine protease (Carrington et al., 1989) defined as a multifunctional protein involved in all essential steps of the potyviral infection cycle (Maia et al., 1996; Syller, 2005), apparently with host specificity (Carbonell et al., 2012; Sænz et al., 2002). Nevertheless, whilst several functions of HCPo may depend on its silencing-suppression activity (Kasschau & Carrington, 2001), the relevance of specific
silencing-suppression mechanisms and the extent of dependence of plant viruses on their natural RSS(s) are largely unknown.

A large number of interactions with viral and host proteins have been attributed to HCPro. Thus, HCPro interacts with the viral coat protein (CP) (Blanc et al., 1997; Roudet-Tavert et al., 2002), cylindrical inclusion (CI) protein (Choi et al., 2000; Guo et al., 2001; Zilian & Maiss, 2011), P1 (Merits et al., 1999), genome-linked protein (VPg) (Guo et al., 2001; Roudet-Tavert et al., 2007; Yambo et al., 2003) and its precursor, nuclear inclusion protein a (Nia) (Guo et al., 2001). As host factor partners, HCPro interacts with two potato RING-finger proteins, HIP1 and HIP2, (Guo et al., 2003), the maize ferredoxin-5 (Cheng et al., 2008), the NtMinD protein (Jin et al., 2007b) and the translation initiation factors eIF(iso)4E and eIF4E (Ala-Poikela et al., 2011) of tobacco, with unknown physiological consequences. HCPro has also been shown to interact with some subunits of the 20S proteasome, inhibiting its endonuclease activity (Ballut et al., 2005; Dielen et al., 2011; Jin et al., 2007a); interactions of HCPro with a calmodulin-related protein (rgs-CaM) (Anandalakshmi et al., 2000) and the ethylene-inducible transcription factor RAV2 (Endres et al., 2010) appear to regulate its silencing-suppression activity. Although these interactions were identified in different heterologous systems, their biological relevance in the context of a natural potyviral infection has not yet been established.

Recent studies with viruses of the family Potyviridae have shown that, even though suppression of silencing is a pivotal potyviral function, a particular infection does not depend on a singular RSS. Hence, plum pox virus (PPV, genus Potyvirus) HCPro can be replaced successfully by the sequence-unrelated RSS P1b from cucumber vein yellowing virus (CVYV, genus Ipomovirus) (Carbonell et al., 2012). Although these two RSSs share no homology in their amino acid sequences, both are present in members of the family Potyviridae and use a similar mechanism, based on siRNA sequestering, to suppress RNA silencing (Valli et al., 2011). To gain further insight into how viruses evade the RNA-silencing machinery of the host, we tested to what extent different RSSs can substitute for HCPro without abolishing PPV infectivity. Our results show that PPV HCPro can be replaced functionally by some, but not all, unrelated RSSs, including the NS1 protein of influenza A virus, and suggest the existence of some specific preferences in the process of counteracting silencing.

**RESULTS**

Generation of a universal PPV intermediate cDNA clone lacking the HCPro silencing suppressor coding sequence

In order to generate PPV-based chimeric viruses easily by replacing HCPro with unrelated proteins, we first generated an intermediate clone in a pGEM-T backbone carrying the PPV P1–P3 cistrons, but lacking the HCPro coding sequence (Fig. 1a). The amplified P1–P3 fragment was engineered to maintain the coding sequence of the two first amino acids of HCPro (SD) just downstream of P1 to ensure efficient P1 self-cleavage. This sequence was followed by an EcoRV/Sbfl cloning site (Fig. 1a). As most proteins aimed to replace the self-cleaving cysteine protease HCPro lack a proteolytic activity to release themselves from the viral polyprotein once infection has taken place, a DNA sequence encoding a PPV Nia protease-cleavage site was added just upstream of the P3 coding sequence (Fig. 1a).

The coding sequence of a set of HCPro-unrelated proteins with suspected RNA silencing-suppression activity [those

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Fig. 1. Schematic representations of viral constructs derived from PPV. (a) Intermediate plasmids created for cloning the coding sequence of different RSSs. P1 and Nia cleavage sites engineered at the ends of the inserted RSSs are also indicated. (b) Chimeric full-length cDNA clones derived from PPV and expressing heterologous RSSs. The GFP reporter gene inserted between the Nb and CP coding sequences is represented by a grey rectangle above the construct.
Several RNA-silencing suppressors support PPV infection

The chimeric virus that carries P1b instead of HCPro has been reported previously (Carbonell et al., 2012), were used as controls. As expected, just a few days post-inoculation (p.i.), GFP foci were detected in all leaves inoculated with PPV and PPV-P1b, whereas no foci appeared in those leaves inoculated with PPV-ΔHC (Fig. 2a; Fig. S1, available in JGV Online). Interestingly, GFP foci also appeared in all leaves inoculated with PPV-P19 (Figs 2a and S1a). The size and number of PPV, PPV-P1b and PPV-P19 GFP foci were similar. Western blot analysis of extracts prepared from GFP focus-containing inoculated tissues showed high accumulation levels of viral CP in plants inoculated with PPV, PPV-P1b and PPV-P19 (Fig. 2b). No virus accumulation was detected in equivalent leaf areas of plants inoculated with PPV-ΔHC (Fig. 2b). Virus-like symptoms (chlorotic mottling) and GFP signals, equivalent to those of PPV and PPV-P1b, were observed in upper non-inoculated leaves of plants infected with PPV-P19 (Figs 2c and S1b). Western blot analysis of extracts prepared from GFP-expressing tissues of systemically infected leaves showed similar virus-accumulation levels for PPV, PPV-P1b and PPV-P19 (Fig. 2d).

A previous report has shown that, whilst the replication of wild-type PPV is maintained actively at late times of infection in N. benthamiana plants, partial recovery occurs in new, growing tissues of plants infected with the chimeric virus carrying P1b instead of HCPro, which is indicative of some defect in the silencing-suppression machinery of the virus (Carbonell et al., 2012). To test the capacity of a PPV chimeric virus with another heterologous RSS to escape from plant recovery, PPV-P19-infected plants were analysed at later stages of the infection process (38 days p.i.) by testing virus accumulation in old and young leaves. As expected, whilst severe symptoms, intense green

TBSV P19 functionally replaces PPV HCPro and efficiently prevents plant recovery

It has been described that PPV HCPro can be replaced functionally by the RSS P1b from the ipomovirus CVYV, another member of the family Potyviridae (Carbonell et al., 2012). P19 is a well-characterized RSS of viruses of the genus Tombusvirus; it suppresses silencing by sequestering double-stranded (ds-)siRNAs (Silhavy et al., 2002; Vargason et al., 2003), the same mechanism that has been proposed for the potyviral HCPro and CVYV P1b (Lakatos et al., 2006; Valli et al., 2011). The infectivity of PPV-P19, in which the P19 coding sequence from tomato bushy stunt virus (TBSV) replaces that of HCPro, was tested by bioassay in Nicotiana benthamiana. Previously described wild-type PPV, HCPro deletion mutant (PPV-ΔHC) (P1ΔHC in Carbonell et al., 2012) and the chimeric virus PPV-P1b, in which HCPro was replaced by CVYV P1b (P1P1b in Carbonell et al., 2012), were used as controls. As expected, just a few days post-inoculation (p.i.), GFP foci were detected in all leaves inoculated with PPV and PPV-P1b, whereas no foci appeared in those leaves inoculated with PPV-ΔHC (Fig. 2a; Fig. S1, available in JGV Online). Interestingly, GFP foci also appeared in all leaves inoculated with PPV-P19 (Figs 2a and S1a). The size and number of PPV, PPV-P1b and PPV-P19 GFP foci were similar. Western blot analysis of extracts prepared from GFP focus-containing inoculated tissues showed high accumulation levels of viral CP in plants inoculated with PPV, PPV-P1b and PPV-P19 (Fig. 2b). No virus accumulation was detected in equivalent leaf areas of plants inoculated with PPV-ΔHC (Fig. 2b). Virus-like symptoms (chlorotic mottling) and GFP signals, equivalent to those of PPV and PPV-P1b, were observed in upper non-inoculated leaves of plants infected with PPV-P19 (Figs 2c and S1b). Western blot analysis of extracts prepared from GFP-expressing tissues of systemically infected leaves showed similar virus-accumulation levels for PPV, PPV-P1b and PPV-P19 (Fig. 2d).

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Table 1. RSSs used to replace PPV HCPro

<table>
<thead>
<tr>
<th>RSS</th>
<th>Virus</th>
<th>Tested by:</th>
<th>Proposed RNA-silencing suppression mechanism</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>P1b*</td>
<td>CVYV</td>
<td>Plants: prevention of silencing in co-agroinfiltration assays, enhancement of PVX pathogenicity</td>
<td>Binding of ds-siRNAs</td>
<td>Valli et al. (2006, 2011)</td>
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<td>P19</td>
<td>TBSV</td>
<td>Plants: prevention of silencing in co-agroinfiltration assays, enhancement of PVX pathogenicity, reversal of transgene silencing, inhibition of VIGS</td>
<td>Binding of ds-siRNAs</td>
<td>Lecellier et al. (2005); Li et al. (2004); Qian et al. (2009); Qiu et al. (2002); Vargason et al. (2003); Voinnet et al. (1999)</td>
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<tr>
<td>NS1</td>
<td>Influenza A virus</td>
<td>Plants: prevention of GFP silencing in co-agroinfiltration assays, enhancement of PVX pathogenicity</td>
<td>Binding of ds-siRNAs and long dsRNAs</td>
<td>Bucher et al. (2004); Delgadillo et al. (2004); Haasnoot et al. (2007); Li et al. (2004)</td>
</tr>
<tr>
<td>P1, P11–383</td>
<td>SPMMV</td>
<td>Plants: prevention of silencing in co-agroinfiltration assays</td>
<td>AGO1 inhibition</td>
<td>Giner et al. (2010)</td>
</tr>
<tr>
<td>P6</td>
<td>CaMV</td>
<td>Plants: reversion of amplicon silencing in transgenic plants</td>
<td>DRB4 inhibition</td>
<td>Love et al. (2007)</td>
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*The chimeric virus that carries P1b instead of HCPro has been reported previously (Carbonell et al., 2012).
fluorescence and high CP levels were maintained in both types of tissue of plants infected with PPV, young leaves of plants infected with PPV-P1b showed a characteristic ‘recovery’ phenotype with the new apical leaves appearing healthy, with much lower intensity of green fluorescence and lower accumulation of CP compared with the older leaves (Fig. 3). However, no recovery was observed in PPV-P19-infected plants, and young leaves still appeared heavily infected at 38 days p.i. (Fig. 3). Intriguingly, whilst the accumulation of wild-type PPV was similar in old and young leaves, a decline in fluorescence intensity and CP levels was apparent in ageing leaves of plants infected with PPV-P19 (Fig. 3).

Thus, our results indicate that CVYV P1b and TBSV P19 can replace HCPro functionally in a PPV infection, but only TBSV P19 prevents the recovery of plants from PPV infection in the absence of HCPro.

**RSSs with different silencing-suppression mechanisms are able to support PPV infection**

Both CVYV P1b and TBSV P19 share with PPV HCPro the ability to bind specifically ds-siRNAs. In order to assess the ability of other proteins that suppress RNA silencing by different mechanisms to replace HCPro in a PPV infection, the HCPro coding sequence of PPV was replaced by sequences encoding the P1 N-terminal region of P1 (P1 N) from sweet potato mild mottle ipomovirus (SPMMV), the P6 from cauliflower mosaic caulimovirus (CaMV) and the protein NS1 from the animal-infecting orthomyxovirus influenza A virus, which respectively suppress RNA silencing by targeting mature RISC complexes (Giner et al., 2010), impairing the DCL4-mediated processing of dsRNAs by interaction with DRB4 (Haas et al., 2008) and an unknown mechanism.

In a new experiment, wild-type PPV and PPV-P19 infected *N. benthamiana* efficiently, as shown previously. Local GFP foci were also detected in six of eight plants inoculated with PPV-NS1, but were smaller in number and size than those of wild-type PPV and PPV-P19 (Fig. 4a and Fig. S2a). No

**Fig. 2.** P19 from TBSV functionally replaces HCPro from PPV. (a) Leaves inoculated with the indicated viruses, taken under an epifluorescence microscope at 9 days p.i. White bar, 1 cm. (b) Western blot analysis of protein extracts prepared from inoculated leaves (two plants per construct) collected at 9 days p.i. (c) Pictures taken under an epifluorescence microscope at 21 days p.i. of the fourth leaves above the inoculated ones. White bar, 1 cm. (d) Western blot analysis of protein extracts prepared from the whole third and fourth leaves above the inoculated ones collected at 21 days p.i. A polyclonal antiserum specific for PPV CP was used for assessment of virus accumulation. Membranes stained with Ponceau red showing RuBisCO are included as loading controls.

**Fig. 3.** A PPV chimera expressing TBSV P19 escapes from plant recovery. (a) Pictures of old and young upper non-inoculated leaves of plants infected with the indicated viruses taken under an epifluorescence microscope at 38 days p.i. White bar, 1 cm. (b) Western blot analysis of protein extracts prepared from upper non-inoculated leaves (two plants per construct) collected at 38 days p.i. O, Old tissues (leaves at positions 6–9 above the inoculated ones); Y, young tissues (leaves at positions 10–13 above the inoculated ones). A polyclonal antiserum specific for PPV CP was used for assessment of virus accumulation. Membranes stained with Ponceau red showing RuBisCO are included as loading controls.
GFP foci were observed in leaves inoculated with PPV-P1N and PPV-P6 (Figs 4a and S2a). Surprisingly, when leaf tissue around the inoculated area was assessed by Western blot analysis, CP accumulation was detected (although at low levels) not only in leaves inoculated with PPV-NS1, but also in those inoculated with PPV-P1N (Fig. 4b). In contrast, no CP accumulation was observed in extracts prepared from plants inoculated with PPV-P6 (Fig. 4b). No disease symptoms were observed in plants inoculated with PPV-NS1, -P1N and -P6. However, some GFP fluorescent spots were detected, with a long delay compared with the wild-type virus, in upper non-inoculated leaves of two plants infected with PPV-NS1 (Figs 4c and S2b). In addition, a faint green fluorescence appeared, with a similar delay to the PPV-NS1 fluorescent spots, in the upper non-inoculated leaves of two plants infected with PPV-P1N (Figs 4c and S2b). Western blot analysis of whole-leaf extracts confirmed the systemic spread of PPV-P1N and PPV-NS1, although the CP accumulation levels of these viruses were very low compared with those of PPV-P19 and PPV wild-type (Fig. 4d). Immunoreactions with available specific antibodies (anti-HP, anti-P19 and anti-NS1) (Fig. 4d) and sequencing of immunocapture (IC)-RT-PCR-amplified products (Fig. S2c) confirmed the identity and the genetic stability of the infecting viruses. Neither GFP fluorescence nor viral CP accumulation was detected in upper leaves of plants infected with PPV-P6 (Fig. 2b, d).

These results indicate that an RSS with a different mechanism of action from those of HCPro, SPMMV P1 and a protein of an animal virus that suppresses silencing by a still-uncharacterized mechanism, the influenza A virus NS1, is able to support a limited systemic infection of a potyvirus in the absence of HCPro. However, not all of the proteins with reported RNA silencing-suppression activity are able to replace HCPro functionally in PPV infection.

**The ability to replace HCPro in the PPV infection process does not correlate strictly with the strength of the silencing suppressor**

To assess the contribution of the RNA silencing-suppression activity of each PPV chimera in its ability to infect *N. benthamiana* plants, each RSS was tested in a co-agroinfiltration assay. Therefore, we constructed *Agrobacterium* binary plasmids expressing the 5′ part of the different chimeric viruses (Fig. 5a), which were co-agroinfiltrated with p35S: GFP (a plasmid expressing single-stranded GFP RNA that is used here as both trigger and reporter of silencing), and pMDC32-NlaPro (a plasmid expressing the protease domain of the PPV Nla protein) (Fig. 5a). For simplicity, in this part of the
In the report, we will refer to each *A. tumefaciens* line by the plasmid that it carries.

Whilst GFP fluorescence declined strongly at 4–5 days post-agroinfiltration (p.a.) in patches expressing p35S:GFP, pMDC32-NlaPro and the pBin19 empty vector, the green fluorescence remained strong at 6 days p.a. in patches co-agroinfiltrated with p35S:GFP and pMDC32-NlaPro plus either pBIN-P1HCPro, pBIN-P1P19, pBIN-P1P1b or pBIN-P1P1N (Fig. 5b), showing that PPV HCPro, TBSV P19, CVYV P1b and influenza A virus NS1 suppress RNA silencing with similar efficiency. In contrast, very weak fluorescence was observed in patches expressing p35S:GFP and pMDC32-NlaPro plus either pBIN-P1P1N or pBIN-P1P6, indicating that, at least in this system, SPMMV P1 N and CaMV P6 have very weak silencing-suppression activity (Fig. 5b).

Northern blot analysis confirmed the green fluorescence observations, showing that GFP mRNA accumulation was similarly high at 6 days p.a. in those leaves expressing TBSV P19, influenza A virus NS1, PPV HCPro and CVYV P1b (Fig. 5c). The drop in GFP mRNA levels could not be prevented in leaves expressing any of the other tested proteins (Fig. 5c).

The genomic expression strategy of potyviruses through proteolytic processing of long polyprotein precursors results in the inevitable presence of extra amino acids, which are necessary for protease recognition in the polyprotein precursor, at the end of the foreign sequences (Fig. 1a). To assess the possibility that these extra amino acids incorporated at the termini of CaMV P6 could be disturbing the silencing-suppression activity of this RSS, then abolishing the infectivity of the PPV-P6 chimera, we compared the anti-silencing activity of wild-type protein with those of their counterparts as engineered in the PPV genome. Hence, we constructed binary plasmids expressing wild-type CaMV P6 and a modified version carrying extra amino acids as they should be produced in the PPV context (SD-P6-QVVVHQ, where SD represents the amino acids +1 and +2 of the P1 cleavage site, whilst QVVVHQ represents the amino acids −6 to −1 of the Nla cleavage site).
Plasmids were co-agroinfiltrated together with p35S: GFP, and their ability to suppress GFP silencing was assessed at 6 days p.a. by monitoring the GFP fluorescence signal (Fig. S3). Intriguingly, wild-type CaMV P6 was unable to suppress RNA silencing in this co-agroinfiltration test, even though it was expressed in the wild-type form, suggesting that a deficiency of P6 anti-silencing activity could be responsible for its inability to support PPV infection.

Altogether, these results indicate that a strong silencing-suppression activity is required for an efficient potyviral infection, but that a weak anti-silencing activity could be enough to support a limited viral infection. However, a strict correlation between HCPro replacement capacity and anti-silencing activity was not observed.

**DISCUSSION**

**How specific is the dependence of potyviruses on HCPro for a successful infection?**

Most plant viruses have been shown to produce RSSs that counteract antiviral defences mediated by RNA silencing (Burgyan & Havelda, 2011; Shimura & Pantaleo, 2011; Valli et al., 2009). However, RSSs are not always essential, and some RSS-defective viruses are able to develop restricted, but still productive, infections (e.g. Ding et al., 1995; Havelda et al., 2003). In turn, HCPro appears to be strictly required for infections caused by viruses of the genus *Potyvirus* (Fig. 2; Carbonell et al., 2012; Garcia-Ruiz et al., 2010). Interestingly, viruses of other genera of the family *Potyviridae* do not depend on HCPro to infect their hosts, as their silencing-suppression activity is provided by other viral proteins (Jansen et al., 2005; Stenger et al., 2005; Valli et al., 2006; Young et al., 2012), and one of these proteins, P1b from the ipomovirus CVYV, can functionally replace the HCPro from the potyvirus PPV (Carbonell et al., 2012). In the present report we show that a heterologous RSS from an unrelated virus, the protein P19 from the tombusvirus TBSV, is also able to support an efficient PPV infection (Fig. 2). PPV-P19 appears to escape antiviral silencing with an efficiency similar to that of the wild-type virus, as plants infected with this chimeric virus do not show a recovery phenotype typical of silencing suppression-deficient viruses (Fig. 3), further supporting the conclusion that potyviruses do not depend on a specific silencing suppression provided by HCPro.

**Do potyviruses depend on a specific RNA silencing-suppression strategy?**

Sequestering of siRNAs appears to be a very successful strategy to suppress silencing, which is used, among other RSSs, by HCPro, P1b and P19 (Lakatos et al., 2006; Valli et al., 2011). However, the lack of similarity and the diverse dependence on specific features at the siRNA ends for efficient binding (Valli et al., 2011) support the idea that siRNA-binding mechanisms of these RSSs are different. The high infectivity of PPV-P1b and PPV-P19 suggests that potyvirus infection does not require a specific strategy of siRNA sequestering and, thus, different mechanisms of siRNA binding could be equally useful for the virus, as long as they provide effective silencing suppression.

To assess whether potyviruses could successfully use other silencing-suppression mechanisms, a series of recombinant viruses in which PPV HCPro was replaced by RSSs with a broad range of anti-silencing strategies was constructed. We were unable to detect infection in the chimeric virus that expressed CaMV P6 (Figs 4 and S2). Recombinant viruses expressing P0 from the polerovirus BWYV, RNase3 from the crinivirus SPCSV, Tat from the retrovirus human immunodeficiency virus (HIV) and RNase XRN4 from *Arabidopsis thaliana*, for which silencing-suppression activities in different experimental systems have been reported (Bennasser et al., 2005; Cuellar et al., 2009; Gazzani et al., 2004), also appear not to be infectious. Unfortunately, we were not able to detect silencing-suppression activity for these proteins when they were expressed as part of the PPV polyprotein in a co-agroinfiltration assay (data not shown). In the case of BWYV P0, and perhaps in other cases, this is due to the extra amino acids added to its ends to facilitate its excision from the viral polyprotein (data not shown). For other proteins, such as CaMV P6, the deficiency could be genuine (Fig. S3). It should be noted that, to our knowledge, for some of these RSSs, such as Tat and P6 viral factors and the *A. thaliana* XRN4 protein, anti-silencing activity using co-agroinfiltration assays has not yet been reported.

Two heterologous RSSs were also able to support PPV infection: the NS1 protein of influenza A virus and an N-terminal fragment of the P1 protein from the ipomovirus SPMMV (Fig. 4). Although NS1 has shown potent RNA silencing-suppression activity in different experimental systems (Table 1; Fig. 5), PPV expressing NS1 instead of HCPro infects *N. benthamiana* very poorly (Fig. 4). NS1 is able to bind long dsRNA and ds-siRNAs, and both capacities are expected to contribute to its ability to suppress the RNA silencing (Bucher et al., 2004; Li et al., 2004). The poor infectivity of PPV-NS1 shows that, even though RNA binding might provide strong silencing-suppression activity, some additional RSS features seem to be required for supporting an efficient potyviral infection.

An N-terminal fragment of the P1 protein from SPMMV (aa 1–383) has been reported to be an efficient RSS (Giner et al., 2010). However, very little silencing-suppression activity was observed here for a similar SPMMV P1 fragment (P1N, aa 1–360) expressed as part of the PPV polyprotein (Fig. 5), which could be due to the extension of this particular deletion or to additional amino acids introduced at both ends of the protein to ensure correct processing during the viral infection. In spite of that, PPV-P1N showed some infectivity (Fig. 4 and Fig. S2), indicating that very little RNA silencing-suppression activity is
enough for a limited potyviral systemic infection. The fact that the infection efficiencies of PPV-NS1 and PPV-P1N are very similar, although the silencing-suppression activity of NS1 is much stronger than that of P1N, suggests that not all anti-silencing mechanisms are equally effective in supporting potyviral infection. P1 has been shown to suppress silencing by Argonaute binding (Table 1), and this strategy appears to be more effective for PPV than the strategy, still to be characterized, used by NS1. It is interesting to remark that, although the infectivity of PPV-NS1 and PPV-P1N chimeric viruses is similar, their patterns of infection are quite different (Fig. 4), suggesting that specific infection features can be conditioned by the RSS used by the virus to counteract the antiviral silencing response of the host.

Only RNA-silencing suppression or something else?

HCPro is a multifunctional protein, known to be involved in all essential steps of viral infection, and it has been shown to interact with a large number of host factors (Ala-Poikela et al., 2011; Anandalakshmi et al., 2000; Ballut et al., 2005; Cheng et al., 2008; Dielen et al., 2011; Endres et al., 2010; Guo et al., 2003; Jin et al., 2007a, b). Some of the functions of HCPro and several of its interactions with host factors appear to be related closely to its RNA silencing-suppression activity (Anandalakshmi et al., 2000; Endres et al., 2010; Kasschau & Carrington, 2001; Kasschau et al., 2003). It is interesting to remark that, whilst the cysteine proteinase activity of HCPro is not necessary for RNA-silencing suppression, it appears to be required for genome amplification (Kasschau & Carrington, 1995), suggesting that potyviral infection could rely on silencing suppression-independent functions of HCPro. Our results do not support this hypothesis as TBSV P19, which lacks any proteolytic activity, is able to provide all HCPro activities essential for PPV infection. We cannot rule out the possibility that this heterologous protein provides a common hypothetical silencing suppression-independent function of HCPro, but, given the large divergence between them, this appears to be very unlikely. What does not seem improbable, however, is that HCPro has important, but not essential, functions in potyviral infection not related to silencing suppression or to its first identified role as helper factor in aphid transmission, which would not be supplied by other heterologous RSSs.

Altogether, our findings demonstrate that the well-characterized RSS HCPro can be replaced by different heterologous RSSs from both plant- and animal-infecting viruses with dissimilar results, thus leading to the conclusion that, whilst PPV infection, and presumably potyviral infections in general, do not rely strictly on a particular RSS, specific silencing-suppression mechanisms would fit better for a given virus. The approach followed here of exchanging the HCPro coding sequence producing heterologous proteins could contribute to identification of novel RSSs and study of their function in the process of a viral infection using a highly ‘silencing-sensitive’ model, such as plants. Moreover, our particular viral system will be very helpful when evaluation of the RNA silencing-suppression activity of a given protein is not feasible in its original viral genome.

METHODS

Plant hosts. Agroinfiltration and viral infectivity assays were performed in N. benthamiana plants. All plants were grown in a greenhouse maintained at 16 h light with supplementary illumination and a temperature range of 19–23 °C.

Plasmids. A detailed description of plasmids used in this study can be found in the Supplementary Methods.

Biolistic inoculation. The Helios Gene Gun system (Bio-Rad) was used for biolistic inoculation. Microcarrier cartridges were prepared from two different clones per construct, with 1.0 µm gold particles coated with the different plasmids at a DNA loading ratio of 2 µg gold mg⁻¹ and a microcarrier loading of 0.5 mg per shot. Helium pressure of 7 bar was used for shooting plants. Each cartridge was shot twice onto two leaves of each plant.

Transient expression by agroinfiltration. N. benthamiana plants were infiltrated with A. tumefaciens C58C1 strain carrying the indicated plasmids as previously described (Valli et al., 2006). Appropriate Agrobacterium cultures were mixed after induction with acetoxyshigine. In the cases of pBIN-P1RSS plasmids expressing RSSs without self-cleaving activity, an Agrobacterium strain carrying a binary plasmid expressing the Nla protease domain (pMDC32-NlaPro) was also included in the infiltration mixtures.

Fluorescence imaging. GFP fluorescence was monitored under long-wavelength UV light (Black Ray model B 100 AP). To capture pictures of fluorescent areas, leaves were examined with a Leica MZ FLIII epifluorescence microscope using excitation and barrier filters at 425/460 nm and 480 nm, respectively, and photographed with an Olympus DP70 digital camera.

Western blot assays. Tissue samples of inoculated leaves were harvested under UV light from GFP-expressing foci or the whole inoculated area, whereas tissue of upper non-inoculated leaves was collected from areas showing green fluorescence or the indicated whole leaves. Control samples corresponding to non-infected leaves were taken from equivalent areas. Preparation of protein samples, SDS-PAGE and electroblotting were done as described previously (Valli et al., 2006). Specific proteins were detected using anti-HCPro rabbit serum, anti-P1b rabbit serum, anti-P19 rabbit serum (kindly provided by Herman Scholthof, Texas A&M University, College Station, TX, USA), anti-NS1 rabbit serum (kindly provided by Ariel Rodriguez, Centro Nacional de Biotecnologia-CSIC, Madrid, Spain) or anti-CP rabbit serum, as primary antibodies, and HRP-conjugated goat anti-rabbit IgG (Jackson) as secondary reagent. The immunostained proteins were visualized by enhanced chemiluminescence detection with a LifeAblot kit (Euroclone). Ponceau red staining was used to check the global protein content of the samples.

IC-RT-PCR. Leaf extracts from infected N. benthamiana plants were homogenized in 5 mM sodium phosphate buffer, pH 7.5 [2 ml (g tissue)⁻¹] and incubated in tubes previously coated with anti-PPV IgGs overnight at 4 °C. The incubation was followed by two washing
steps with PBS–TWEEN buffer (16 mM PBS, 0.1 M NaCl, 0.5 g Tween 20 1–1, pH 7.2). RT-PCR was performed using a Titan kit (Roche Molecular Biochemicals) with primers targeting the 3’ end of PPV P1 (#90: 5’-CGGACCCAATGCGAG-3’) and 5’ end of PPV P3 (#317: 5’-TGAACCTACTTTGACAG-3’). For further sequence analysis, PCR fragments were purified using a MinElute PCR purification kit (Qiagen).

**RNA extraction and Northern blot analysis.** Samples containing mRNAs were prepared from agroinfiltrated leaf tissue and subjected to Northern blot analysis as described previously (Valli et al., 2006).

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