Tobacco etch virus P1 protein traffics to the nucleolus and associates with the host 60S ribosomal subunits during infection

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Genus *Potyvirus* comprises a large group of positive-strand RNA plant viruses whose genome encodes a large polyprotein processed by three viral proteinases. P1 protein, the most amino-terminal product of the polyprotein, is an accessory factor stimulating viral genome amplification whose role during infection is not well understood. We infected plants with *Tobacco etch virus* (TEV; genus *Potyvirus*) clones in which P1 was tagged with a fluorescent protein to track its expression and subcellular localization or with an affinity tag to identify host proteins involved in complexes in which P1 also takes part during infection. Our results showed that TEV P1 exclusively accumulates in infected cells at an early stage of infection, and that the protein displays a dynamic subcellular localization, trafficking in and out of the nucleus and nucleolus during infection. Inside the nucleolus, P1 particularly targets the dense granular component. Consistently, we found functional nucleolar localization and nuclear export signals in TEV P1 sequence. Our results also indicated that TEV P1 physically interacts with the host 80S cytoplasmic ribosomes and specifically binds to the 60S ribosomal subunits during infection. *In vitro* translation assays of reporter proteins suggested that TEV P1 stimulates protein translation, particularly when driven from the TEV internal ribosome entry site. These *in vitro* assays also suggested that TEV helper-component proteinase (HC-Pro) inhibits protein translation. Based on these findings, we propose that TEV P1 stimulates translation of viral proteins in infected cells.

In this work, we researched the role during infection of tobacco etch virus P1 protease. P1 is the most mysterious protein of potyviruses, a relevant group of RNA viruses infecting plants. Our experiments showed that the viral P1 protein exclusively accumulates in infected cells at an early stage of infection and moves in and out of the nucleus of infected cells, particularly targeting the nucleolus. Our experiments also showed that P1 protein binds host ribosomes during infection. Based on these findings and other *in vitro* experiments we propose that P1 protein stimulates translation of viral proteins during infection.
Plant viruses have evolved as a combination of genes encoding proteins displaying a limited series of catalytic activities but with the ability to interact with specific cellular factors to successfully achieve virus genome replication and gene expression, virion assembly, virus movement, and to counter host defense systems. **Tobacco etch virus** (TEV) is a member of the genus **Potyvirus** (one of the nine currently accepted genera within the family **Potyviridae**) which by itself includes approximately 30% of known species of plant viruses. The genome of potyviruses consists of a single-stranded, positive-sense RNA of approximately 10,000 nucleotides (nt) that is linked at the 5’ end to a viral protein genome-linked (VPg) and contains a poly(A) tail at the 3’ end. This genomic RNA includes a long open reading frame encoding a polyprotein that, when translated, is cleaved by three viral proteases into at least ten mature proteins: P1, helper-component proteinase (HC-Pro), P3, 6K1, cylindrical inclusion (CI), 6K2, nuclear inclusion a (Nla; a polypeptide which is further processed to produce VPg and NlaPro), nuclear inclusion b (Nlb) and the coat protein (CP) (1).

Additionally, the protein P3N-PIPO is produced through a translational frame shift in the P3 cistron (2).

Most potyviral proteins have been attributed important roles during the infectious cycle (3). HC-Pro, a cysteine proteinase that self-cleaves from the viral polyprotein, suppresses the host defensive RNA silencing pathways (4). It is also involved in aphid transmission (5). P3N-PIPO and CI play roles in virus cell-to-cell movement (6). 6K2 anchors the viral replication complex to intracellular membranes (7, 8). VPg acts as a primer during viral RNA synthesis (9), establishes crucial interaction with host factor eIF4E or eIF(iso)4E (10), and is involved in translation (11). NlaPro is a serine protease that in **cis** and in **trans** cleaves most of the potyviral polyprotein (12). Nla is also involved in virus replication (13). Nlb is the viral RNA-dependent RNA polymerase (14) and CP, in addition to virion assembly, is involved in aphid transmission (15), and viral movement (16). In contrast, P1, the first product of potyviral polyprotein, is a rather mysterious protein whose functions during the infectious cycle are mostly unknown (17). The P1 cistron is the most variable in size and sequence in the various species of the genus (18). The P1 protein contains a carboxy-terminal serine protease domain that catalyzes its own cleavage from the viral polyprotein (19), and exhibits a strong RNA binding activity (20, 21). Although P1 contributes to the virus infectious cycle, it is not essential, because a virus deletion mutant completely lacking the P1 cistron, although debilitated, is still viable (22, 23). Although P1 protein enhances viral suppression of RNA silencing mediated by HC-Pro (24, 25), it has been recently shown that more HC-Pro accumulates if HC-Pro is translated as P1/HC-Pro fusion than alone (26).
However, it was recently shown that P1 protein is also important in defining virus host range (18, 27). In this work we investigated the roles of P1 protein in TEV infection. Towards this end, we analyzed P1 expression and subcellular localization during infection. We also identified host proteins that form complexes with P1 during infection. We found that P1 accumulates in infected cells at an early stage of the infectious process and then mostly disappears. We also found that P1 initially localizes in the nucleolus and then traffics back to the cytoplasm. We demonstrated that TEV P1 contains a functional nucleolar localization signal (NoLS) and a nuclear export signal (NES). In infected plant tissues, we encountered P1 mostly associated with ribosomal proteins. We demonstrated that P1 binds in vivo to host 80S cytoplasmic ribosomes, and more specifically to the 60S ribosome subunits. Based on this interaction, we investigated a possible role of P1 in translation and found that, in a wheat germ system, P1 stimulates translation of reporter proteins in vitro, particularly when translation is driven from the TEV internal ribosome entry site (IRES). Based on these findings, we propose a model in which potyviral P1 protein binds 60S ribosomal subunits to subvert the host translation machinery during infection.

MATERIALS AND METHODS

Recombinant TEV clones and plant inoculation. Various recombinant TEV clones were constructed using common molecular biology techniques and starting from pGTEVa (28), a binary plasmid with a cassette to agroinoculate wild-type TEV (TEV-wt; GenBank accession number DQ986288 including silent mutations G273A and A1119G) under the control of Cauliflower mosaic virus (CaMV) 35S promoter and terminator. TEV-VenusP1 contained the fluorescent protein Venus, a green-yellow derivative of Aequorea victoria green fluorescent protein (GFP) (29), fused to the amino terminus of P1 (Fig. 1A). TEV-GFP contained the enhanced GFP between the P1 and HC-Pro cistrons (Fig. 1A). In this clone, the GFP is released from the viral polyprotein through the proteolytic activities of P1 and NlaPro proteinases (28). In TEV-VenusP1-mCherryNlb, in addition to the Venus fusion to P1, the red fluorescent protein mCherry (30) was fused to the amino terminus of Nlb (Fig. 1A). Finally, in TEV-TSTP1, the Twin-Strep-tag (TST), commonly used for protein purification by affinity chromatography (31), was fused to the amino terminus of P1 (Fig. 1A). Sequencing of the corresponding plasmids confirmed the correct construction of all these recombinant TEV clones. Their exact sequences are specified in the supplemental material (Fig. S1). Nicotiana benthamiana Domin plants were kept in a growth chamber with a photoperiod of 12 h light at
25 °C and 12 h dark at 23 °C. Plant agroinoculation was performed using *Agrobacterium tumefaciens* C58C1 (harboring the helper plasmid pCLEAN-S48) at optical density of 0.5 at 600 nm as described (32). TEV virions were partially purified from infected tissues and used for mechanical inoculation of *N. benthamiana* leaves (32).

**Protein transient expression in *N. benthamiana* leaves.** For transient expression, leaves of *N. benthamiana* plants were infiltrated with cultures of *A. tumefaciens* C58C1 (harboring the helper plasmid pCLEAN-S48) as described (33). To express *Arabidopsis thaliana* fibrillarin 2 (AtFib2) (34, 35), fused to a red fluorescent protein (AtFib2mRFP), *A. tumefaciens* was transformed with a binary plasmid previously described (35). To express *A. thaliana* ribosomal protein L24B (AtRPL24B) (36), fused to mCherry (AtRPL24BmCherry), the AtRPL24B cDNA was amplified from *A. thaliana* Col-0 by RT-PCR and cloned in a modified version of pEarlyGate101 (Invitrogen). To express Venus, VenusP1 and derivatives of Venus P1, *A. tumefaciens* was transformed with a series of binary plasmids based on a modified version (32) of pCLEAN-G181 (GenBank accession number EU186083) containing, between the left and right border of the T-DNA, expression cassettes consisting of CaMV 35S promoter, a modified version of *Cowpea mosaic virus* (CPMV) RNA-2 5’ untranslated region (UTR) (37), the cDNA of the corresponding protein, CPMV RNA-2 3’ UTR and CaMV 35S terminator. The exact sequence of each construct is specified in the supplemental material (Fig. S2).

**Analysis of fluorescent proteins.** Expression of fluorescent proteins was analyzed by confocal laser scanning microscopy using a Leica TCS SL with a HCX PL APO 40X/1.25-0.75 oil lens. Venus and mCherry were detected with excitation lasers of 488 and 543 nm, and detection windows of 520-550 and 610-670 nm, respectively. Optical section was 1 μm. Also, with a fluorescence stereomicroscope Leica MZ 16 F equipped with filters DSR and GFP2 (Leica). Fluorescent infection foci were analyzed using ImageJ software.

**Infectivity assays.** TEV recombinant clones containing mutations in the P1 cistron (see supplemental material; Fig. S3) and the transcription factor Rosea1 (Ros1) as a reporter marker (28) were agroinoculated in two different leaves of batches of 10 *N. benthamiana* plants. Infection symptoms in these plants were recorded over time by visual inspection. Viral load was estimated from the anthocyanin accumulation induced by the Ros1 reporter marker activity. Anthocyanins in infected tissues from three different plants for each viral construct were extracted in acidified methanol and quantified spectrophotometrically at 530 nm (28).

**Protein analysis.** Proteins were separated by denaturing (0.5% sodium dodecyl sulfate; SDS) polyacrylamide gel electrophoresis (PAGE) in 12.5% polyacrylamide gels,
electroblotted to PVDF membranes (GE Healthcare) and analyzed by western blotting as described (38). A monoclonal antibody against the TST tag (StrepMAB-classic horseradish peroxidase, IBA) was used at a 1:5000 dilution and a polyclonal antibody against the TEV CP (conjugated to alkaline phosphatase, Agdia) at a 1:10,000 dilution. Immunoblots were quantified with a luminescent image analyzer (LAS-3000, Fujifilm) as described (28).

Polysomes were isolated from 2 g of TEV-TSTP1-infected *N. benthamiana* plants, harvested at 5 days post-inoculation (dpi), as described (39). Tissue was ground in a mortar in the presence of liquid N$_2$ and homogenized with 10 volumes extraction buffer (200 mM Tris-HCl, pH 9.0, 400 mM KCl, 200 mM sucrose, 35 mM MgCl$_2$, 5 mM dithiothreitol (DTT), 30 mM ethylenediaminetetraacetic acid (EDTA) and a cocktail of protease inhibitors (Complete, Roche Applied Sciences). Extract was clarified by centrifugation at 25,000 x g for 10 min.

Five ml aliquots of the supernatants were centrifuged through 4 ml layers of 1.75 M sucrose in 40 mM Tris-HCl, pH 9.0, 200 mM KCl, 30 mM MgCl$_2$, 5 mM DTT and either 30 mM or 5 mM EDTA to separate or not ribosomal particles, and centrifuged at 340,000 x g for 1 h. Sediments were resuspended in a total of 1 ml of 40 mM Tris-HCl, pH 8.5, 200 mM KCl, 30 mM MgCl$_2$, 5 mM DTT and either 30 mM or 5 mM EDTA, and layered on continuous 15-60% sucrose gradients in 40 mM Tris-HCl, pH 8.5, 20 mM KCl, 10 mM MgCl$_2$, 5 mM DTT, 10 mM EDTA. Tubes were centrifuged for 3 h at 4 °C in a SW 40 Ti rotor (Beckman Coulter) at 40,000 rpm (285,000 x g) and 16 fractions collected.

**Purification of protein complexes containing TEV TSTP1 and protein identification by mass spectrometry analysis.** Symptomatic *N. benthamiana* leaf tissues (15 g) infected by TEV-wt or TEV-TSTP1 (Fig. 1A) were harvested at 5 dpi, ground in a mortar with liquid N$_2$, and homogenized with 45 ml of extraction buffer (100 mM Tris-HCl, pH 8.0, 150 mM KCl, 10 mM MgCl$_2$, 10 mM DTT, 1 mM EDTA, 1% Nonidet P-40) containing a cocktail of protease inhibitors (Complete, Roche). The crude extracts were clarified twice by centrifugation at 4 °C, first at 12,000 x g for 15 min and then at 95,000 x g for 30 min. The TST-tagged P1 protein was purified by chromatography using a 1-ml Strep-tactin Superflow column (IBA) with an ÄKTA Prime Plus liquid chromatography system (GE Healthcare) operated at 4 °C at a flow rate of 1 ml/min. After equilibration with 10 ml of extraction buffer, the column was loaded with the clarified extract and washed with 20 ml of extraction buffer.

Bound protein complexes were eluted with 20 ml of extraction buffer containing 10 mM D-desphiothietin, collecting 0.5-ml fractions. Fractions from the TEV-TSTP1 infected tissues were analyzed by western blot with the anti-TST antibody and those containing substantial amounts of TSTP1, pooled and the protein precipitated adding 4 volumes of 12.5%
trichloroacetic acid, 10 mM DTT in acetone. The same process was followed with the corresponding fractions eluted in the control purification process from tissues infected by TEV-wt. Protein preparations were separated by SDS-PAGE (12.5% polyacrylamide, 0.05% SDS) and the gel stained with Coomassie Blue. Whole lanes corresponding to each sample were excised from the gel, cut in pieces, and the proteins subjected to in-gel digestion with sequencing grade trypsin (Promega) as described (40). Peptides were eluted from the gel pieces and analyzed by liquid chromatography and tandem mass spectrometry as previously described (33).

**In vitro translation assays.** The TNT Coupled Wheat Germ Extract System (Promega) was used to translate proteins *in vitro* in the presence of L-[35S]-methionine (1000 Ci/mmol). Reactions were started for 15 min at 30 °C with 0.2 pmol of a series of plasmids with cassettes to translate the firefly luciferase (Fluc), TEV truncated P1, P1, HC-Pro and P1/HC-Pro. Then, 0.2 pmol of plasmids with monocistronic or bicistronic reporter cassettes were added to all reactions and incubation continued for 40 min (monocistronic reporter) or 2 h 15 min (bicistronic reporter). The exact sequences of the cassettes subjected to *in vitro* translation are in supplemental material (Fig. S4). Translation products were separated by SDS-PAGE (12.5% polyacrylamide, 0.05% SDS), the gel fixed in 20% methanol and 10% acetic acid for 30 min, dried under vacuum and analyzed by phosphorimetry (Fujifilm FLA-5100).

**RESULTS**

**P1 expression during TEV infectious cycle.** To learn about the roles of P1 protein in the TEV infectious cycle, we infected plants with a recombinant TEV clone in which the P1 protein was tagged with a fluorescent protein to track its expression and subcellular localization. We inserted a cDNA coding for Venus, as an amino terminal fusion to P1 in an infectious TEV clone, obtaining TEV-VenusP1 (Fig. 1A). We agroinoculated *N. benthamiana* plants with TEV-VenusP1 and, as controls, with TEV-wt and TEV-GFP (Fig. 1A). Plants agroinoculated with TEV-VenusP1 became infected showing mild disease symptoms with a one-day delay with respect to TEV-wt and TEV-GFP (Fig. 2A). However, since P1 is a non-essential protein for the virus, it is difficult to evaluate the effect of the tag on P1 function.

Analysis with a fluorescence stereomicroscope of symptomatic non-inoculated (systemic) leaves of plants infected by TEV-VenusP1 showed faint green fluorescence exclusively at the infection front. In contrast, systemic leaves of plants infected by TEV-GFP displayed a strong green fluorescence in all symptomatic tissues. We checked the stability of
the TEV-VenusP1 clone by reverse transcription (RT) polymerase chain reaction (PCR) amplification of viral cDNAs from RNA preparations obtained from infected tissues (Fig. 2B). Sequence analysis confirmed the presence and correct insertion of the Venus cDNA in the genome of the viral progeny. Furthermore, new plants infected with virions purified from plants initially agroinoculated with TEV-VenusP1 exhibited the same pattern of green fluorescence in the periphery of symptomatic areas. In contrast to agroinoculation, mechanical inoculation of purified virions allows analysis of infection foci in inoculated leaves. Whereas infection foci of TEV-GFP appeared uniformly fluorescent when observed under a fluorescence stereomicroscope, infection foci of TEV-Venus P1 exhibited only a green fluorescent ring at the periphery (Fig. 1B).

We constructed a new recombinant TEV clone in which, in addition to the Venus fusion to P1, we also tagged Nb with the red fluorescent protein mCherry. In this new clone (TEV-VenusP1-mCherryNb; Fig. 1A), mCherry was inserted as an amino-terminal fusion to Nb. Agroinoculation of *N. benthamiana* plants with TEV-VenusP1-mCherryNb demonstrated that this recombinant clone was also infectious, although mild disease symptoms appeared with a three-day delay with respect to TEV-wt (Fig. 2). This clone allowed observation of the green fluorescence of VenusP1 and the red fluorescence of mCherryNb in the same infection foci. Whereas red fluorescence was detected throughout the foci, green fluorescence was only observed in the peripheral rings (Fig. 1C). Taken together, these results suggest a transient accumulation of the fusion protein VenusP1 at an early stage of viral infection. Since potyviruses express their proteins (except for P3N-PIPO) from a single large polyprotein, potyvirus proteins are synthesized in equimolar amounts and relative changes in accumulation must result from different degradation rates. Therefore, these results suggest that TEV P1, in contrast to other viral proteins, reaches the highest level in the infected cells at an early stage of infection, and then at late stages, P1 is efficiently degraded.

To obtain further support for the above hypothesis, we constructed a new recombinant TEV clone in which P1 was tagged at the amino terminus with TST. The new recombinant clone (TEV-TSTP1; Fig. 1A) was agroinoculated into *N. benthamiana* plants alongside plants inoculated with TEV-wt. TEV-TSTP1 was infectious and agroinoculated plants displayed, again, mild symptoms with a one-day delay with respect to TEV-wt (Fig. 2). In plants infected by TEV-TSTP1, the third leaf above the agroinoculated one was harvested in triplicate on various dpi. Proteins were extracted and the accumulation of viral TSTP1 and CP determined by SDS-PAGE followed by western blot analysis with anti-TST and anti-CP antibodies. A representative western blot corresponding to a time-course of one of the three
sample replicates is shown in Figure 1D. Figure 1E shows the dynamics of TSTP1 and CP accumulation in systemic leaves of infected plants. Whereas TEV CP accumulates continuously in the infected tissue from 4 to 14 dpi, TSTP1 increases accumulation from 4 to 6 dpi and then quickly falls. This result supports a transient accumulation of P1 during the early stage of infection and subsequent effective degradation at later stages.

P1 subcellular localization in infected cells. To investigate P1 subcellular localization during infection, we used the infectious TEV-VenusP1-mCherryNIb clone (Fig. 1A). NIb subcellular localization is well known (41, 42), and served as an internal control in these experiments. *N. benthamiana* plants were mechanically inoculated with TEV-VenusP1-mCherryNIb virions and infection foci were analyzed by confocal laser scanning microscopy. Green and red fluorescence reported VenusP1 and mCherryNIb subcellular localization, respectively. We analyzed the accumulation of both fluorescent proteins from the periphery to the center of infection foci (Fig. 3A). Localization of VenusP1 differed across the infection foci (Fig. 3B). As already mentioned, green fluorescence was brightest at the periphery of the infection foci, just at the viral replication front. In these cells, VenusP1 was mostly localized in a large subnuclear body, presumably the nucleolus, and also more diffusely in the nucleus and cytoplasm (Fig. 3B, I). mCherryNIb was still not detectable in these cells (Fig. 3B, I). In adjacent cells towards the center of the infection foci, VenusP1 was again detected in nucleolus, nucleus and cytoplasm, whereas mCherryNIb was detected in the nucleus and cytoplasm (Fig. 3B, II). The intensity of green fluorescence progressively decreased in cells towards the epicenter of the infection foci. Nonetheless, it was still possible to observe VenusP1 in the nucleus and cytoplasm but no longer in the nucleolus (Fig. 3B, III and IV). Inside the nucleus, VenusP1 displayed a more perinuclear localization than mCherryNIb (Fig. 3B, II to IV). At the center of the infection foci, VenusP1 was no longer detected (Fig. 3B, V). mCherryNIb was always detected in the nucleus and cytoplasm, as expected (Fig. 3B, II to V) (41, 42). These results indicate that TEV P1 displays a dynamic intracellular localization during the cellular infectious cycle. At the beginning of the infection, the protein localizes in the cytoplasm and nucleus, particularly targeting the nucleolus. Then, the protein seems to exit the nucleolus.

To confirm and further investigate P1 nucleolar localization, we performed co-localization experiments between VenusP1 and well known nucleolar proteins: AtFib2 (34, 35) and AtRPL24B (36). AtFib2 and AtRPL24B were transiently expressed in *N. benthamiana* leaves by means of infiltration with *A. tumefaciens* cultures. Infiltrated leaves were mechanically inoculated 6 h early with TEV-VenusP1 virions. AtFib2 and AtRPL24B...
were expressed as amino terminal fusions to monomeric red fluorescent protein (mRFP; AtFib2mRFP) and mCherry (AtRPL24BmCherry), respectively. Whereas AtFib2 localizes to the nucleolus dense fibrillar component and Cajal bodies, AtRPL24B localizes to nucleolus dense fibrillar and granular components, as well as the nucleus and cytoplasm. Three days after the inoculation and subsequent infiltration, we could observe under the confocal microscope both the red fluorescence of the transiently expressed proteins in the whole agroinfiltrated area and the green fluorescence of the viral VenusP1 at the periphery of the infection foci. Nucleolar localization of marker AtFib2mRFP and AtRPL24BmCherry proteins in infected cells was the same as in uninfected cells. The intense green fluorescence of VenusP1 co-localized with the red fluorescence of AtFib2mRFP in the large subnuclear body, confirming that this actually was the nucleolus (Fig. 4A). However, the merged image with the two fluorescence signals indicated very little intranucleolar co-localization (Fig. 4A). Comparison with AtFib2mRFP showed that VenusP1 was also absent from the Cajal bodies (Fig. 4A). In contrast, the merged image indicated a better co-localization between VenusP1 and AtRPL24BmCherry at the periphery of the nucleolus (Fig. 4B), corresponding to the granular component (Fig. 4C). These results support the nucleolar localization of TEV P1 during the early stage of viral infection; more specifically our results suggest that the protein traffics to the nucleolus granular component.

Nucleolar localization and nuclear export signals in P1 sequence. The results shown above about P1 subcellular localization during the TEV infectious cycle suggest that this protein may traffic from the cytoplasm to the nucleus, particularly targeting the nucleolus, and then back to cytoplasm, like the ribosomal protein AtRPL24B. We searched for putative NoLS and NES in TEV P1 sequence using the NoD (43) and NetNES (44) algorithms, respectively. NoD predicted a single NoLS between TEV P1 amino acids 89 and 109 (LTHGKRRKVSVNNKRNRRRKV; Fig. 5). NetNES predicted a single NES between amino acids 251 and 260 (LTFGSSGLVL; Fig. 5). We tested the functionality of these motifs by A. tumefaciens-mediated transient expression in N. benthamiana leaves of a series of VenusP1 constructs containing deletions and amino acid substitutions in the P1 coding sequence (Fig. 6A and Fig. S2).

When wild-type VenusP1 (Fig. 6A, I) was transiently expressed in N. benthamiana leaves, analysis by confocal laser scanning microscopy of the infiltrated tissues 3 days post-infiltration showed an intense green fluorescence in the nucleolus. This fluorescence was less intense in the nucleus and an even less in the cytoplasm (Fig. 6B, I). In contrast, transient expression of a control construct consisting of Venus alone (Fig. 6A, II) showed similar green
fluorescence intensity in both nucleus and cytoplasm, but largely excluding the nucleolus (Fig. 6B, II). We split P1 in two halves: the amino-terminal half from amino acid 1 to 115 containing the presumed NoLS and the carboxy-terminal half from amino acid 116 to 304 (Fig 6A, III and IV). Transient expression of each half fused to the carboxyl terminus of Venus showed green fluorescence in the nucleolus exclusively in the case of the construct containing amino acids 1-115 (Fig. 6B, compare III and IV). Therefore, the first 115 amino acids of P1 are necessary and sufficient for its nucleolar localization. Next, we fused the 21 amino acids of the putative NoLS (from P1 amino acid 89 to 109) to the amino terminus of Venus (Fig. 6A, V). This polypeptide was sufficient to transport Venus inside the nucleolus (Fig. 6B, V), indicating that it contains a functional NoLS. This sequence contains two stretches of basic amino acids: 93-KRRK-96 and 102-KRNRRRK-108, the second interrupted by Asn-104. We mutated wild-type VenusP1 to obtain three different constructs. In the first, each amino acid of the motif 93-KRRK-96 was mutated to Ala (Fig. 6A, VII). In the second, each basic amino acid of the motif 102-KRNRRRK-108 was mutated to Ala (Fig. 6A, VIII). In the third, both mutated motifs were combined (Fig. 6A, IX). No mutant reached the nucleolus (Fig. 6B, VII to IX). Taken together, these results demonstrated the existence of a NoLS in TEV P1 constituted by two motifs of basic amino acids, both of them required for entering the nucleolus.

Next, we deleted the putative NES (251-LTFGSSGLVL-260) from the P1 sequence in the VenusP1 construct (Fig. 6A, VI). Inspection of the infiltrated leaves showed green fluorescence in the nucleus and nucleolus; but in contrast to wild-type VenusP1, no fluorescence was detected in the cytoplasm (Fig. 6B, compare I and VI). We, next, mutated the three carboxy-terminal amino acids of the presumed NES (258-LVL-260) to three consecutive Ala (Fig. 6A, X) in the VenusP1 construct. Again, green fluorescence was detected in nucleus and nucleolus but not in the cytoplasm. These results support the notion that TEV P1 contains a functional NES involved in the protein export from the nucleus to the cytoplasm.

Effect of P1 mutations on viral infection. We tested the effect of some of the P1 mutations described above on TEV infectivity and viral load. To this aim, we introduced a series of mutations in the P1 cistron of a TEV recombinant clone that contains the transcription factor Ros1 as a reporter marker (28). The resulting mutant viral clones (Fig. S3), along with the control including the wild-type P1, were agroinoculated in batches of N. benthamiana plants. The number of infected plants was recorded over time for the different mutants. Viral load was estimated by measuring the anthocyanin accumulation induced by the
Ros1 reporter at 3 days after symptoms first emerged in each infected plant. In TEV clones tagged with Ros1, anthocyanin accumulation correlates with viral load (28). As already reported (22, 23), the whole deletion of the P1 cistron in the TEV genome renders a viable virus, although reduces infectivity and viral load with respect to wild-type virus (mutant Mt-ΔP1, Fig. 7A and B). This is in contrast to what occurs with mutation of the His 214 to Ala in P1 catalytic triad of the serine protease domain that abolishes infectivity (mutant Mt-H214A, Fig. 7A). Also as reported (22, 23), this defect was partially rescued by inserting a subrogate NlaPro cleavage site (mutant Mt-H214+NlaPro, Fig. 7A and B). Mutation of the LVL motif in the P1 NES (mutant X in Fig. 6) rendered a non-infectious virus (Mt-X, Fig. 7A). As this motif is present in the serine protease domain, we also assayed this mutant but inserting a subrogate NlaPro cleavage site (Mt-X+NlaPro) to distinguish the nuclear export and the proteolytic functions. In this case, infectivity and viral load were recovered, but only in part (Mt-X+NlaPro, Fig. 7A and B). Mutation of the basic amino acids in the P1 NoLS (mutants VII, VIII and IX in Fig. 6) had little effect on TEV infectivity, but reduced viral load in infected plants (Mt-VII, Mt-VIII and Mt-IX, Fig. 7A and B).

Host proteins associated to P1 during TEV infection. To further investigate the roles of P1 during TEV infection, we wished to identify host proteins involved in complexes in which P1 is also a component during infection. For this purpose, we inoculated *N. benthamiana* plants with the infectious TEV-TSTP1 clone (Fig. 1A). Control plants were also inoculated with TEV-wt. Symptomatic tissues were harvested 5 dpi (an early stage of infection when P1 concentration in infected tissue is high, see Fig. 1E) and homogenized. The tagged P1 (TSTP1) was purified from the plant extract by liquid chromatography under native conditions using a Strep-tactin column. This tag has previously been used to successfully purify complexes involving potyviral proteins from infected tissues (45). The same purification process was applied to an extract from tissues infected by TEV-wt, as a negative control. Electrophoretic analysis of the chromatographic fractions showed enrichment of a number of proteins from the tissues infected by TEV-TSTP1 relative to those infected by TEV-wt (Fig. 8A, compare lanes 2 and 3). Moreover, the successful purification of TSTP1 was validated by a western blot analysis using an anti-TST antibody (Fig. 8A, lower panel). Purified proteins were digested in gel with trypsin and the resulting peptides were extracted and characterized by liquid chromatography separation followed by tandem mass spectrometry analysis. To obtain a comprehensive list of host proteins actually associated to TEV P1 during infection and to eliminate possible false interactors, this analysis was performed independently in two different samples purified from two different batches of
tissues infected by TEV-TSTP1 and in one sample purified from tissues infected by TEV-wt (Table S1). Table 1 shows the refined list of 32 protein hits common to both TEV-TSTP1 infected samples and absent from the TEV-wt infected control. Surprisingly, most of the hits corresponded to ribosomal proteins, particularly components of the cytosolic 60S ribosomal subunit (15 proteins). The list also contained two proteins components of the 40S ribosomal subunit, one protein involved in translation and four chaperones. In addition, TEV P1, HC-Pro and CP were also identified. These results suggest that TEV P1 interacts with the plant ribosome during infection.

To test this hypothesis, we purified polysomes from N. benthamiana tissues infected by TEV-TSTP1. The polysome preparation was either treated or not with EDTA to separate the ribosomal subunits and subsequently centrifuged on sucrose gradients (39, 46). After centrifugation, the gradients were fractionated and the optical density at 254 nm of the different fractions measured. The presence of the two large ribosomal RNAs (18S and 28S) in the gradient fractions was verified by an electrophoretic analysis in an agarose gel followed by ethidium bromide staining. Finally, the presence of TSTP1 in the fractions was examined by SDS-PAGE and western blot analysis. When the polysomes preparation was not treated with EDTA, TSTP1 was mainly detected in the fractions containing the complete 80S ribosomes (Fig. 8B). When the preparation was treated with EDTA, TSTP1 was exclusively detected in the fractions containing the 60S ribosomal subunit (Fig. 8C). These results support the physical association of TEV P1 with the large subunit of cytosolic ribosomes in actively-translating ribosomes during the infectious process.

We investigated whether the nucleolar localization of P1 is required for binding the host 60S ribosome subunits. To this aim we inserted the mutations that abolish nucleolar localization of TEV P1 (Mt-IX; Fig. 6A and Fig. 6B, IX) into the TEV-TSTP1 infectious clone and infected plants. Ribosomal 60S subunits were purified from infected tissue at 5 dpi by sucrose gradient centrifugation. SDS-PAGE and western blot analysis of the fractions showed that P1 Mt-IX is still able to specifically bind the ribosomal 60S subunits (Fig. 8D), indicating that lack of nucleolar localization does not preclude P1 to bind the host 60S particles.

**Effect of P1 on protein translation in vitro.** The results shown above may suggest an unexpected role of TEV P1 in protein translation during the infectious cycle. To get some insight into this hypothesis, we performed in vitro translation assays. We tried to produce recombinant TEV P1 in Escherichia coli to study its effect on in vitro translation, but the expressed protein accumulated in inclusion bodies and we were unable to purify it as a
soluble form in native conditions. For this reason, we chose a coupled *in vitro* transcription-translation system (47, 48), to co-express TEV P1 or control proteins with reporter constructs to search for a possible effect of P1 on translation. *In vitro* translations were carried out using the wheat germ extract containing bacteriophage SP6 RNA polymerase to couple *in vitro* transcription and translation of the desired cDNAs, and [35S]methionine to track protein synthesis. The *in vitro* translation extract was programmed with reporter plasmids containing the bacteriophage SP6 promoter and a poly(A) tail, designed to express the fluorescent protein Venus, tagged at the amino terminus with the human influenza virus haemaglutinin tag, flanked with a modified version of CPMV RNA-2 5’ and 3’ UTRs (construct I, Fig. 9A and Fig. S4). We used CPMV RNA-2 5’ and 3’ UTRs because these elements very efficiently promote translation of heterologous proteins in plants, particularly the modified version of the 5’ UTR used here that contains a specific mutation eliminating an internal ATG codon (37).

Fifteen min ahead of adding the reporter plasmid, aliquots of the wheat germ extract were programmed with normalized amounts of plasmids containing expression cassettes to produce: (1) Fluc, taken as a negative control; (2) a truncated version of TEV P1 (155 initial amino acids); (3) wild-type TEV P1; (4) wild-type TEV HC-Pro—the second protein in TEV polyprotein, though not being involved in translation, was initially selected as a negative control—and (5) a polyprotein consisting of TEV P1/HC-Pro, such as they appear in the viral genome. These expression cassettes consisted of the SP6 phage promoter, modified CPMV RNA-2 5’ UTR, the cDNAs coding for the corresponding protein, and CPMV RNA-2 3’ UTR followed by a poly(A) tail (constructs III to VII, Fig. 9A and Fig. S4). Reactions were allowed to continue for 40 more min. A prospective time-course experiment showed that *in vitro* translation reactions were not saturated at this point (data not shown).

Three independent replicates of the experiments were performed. Translation products were separated by SDS-PAGE and quantified by phosphorimager analysis. Figures 9B and C show the effect of the expressed proteins on translation. Figure 9D shows the autoradiogram corresponding to one of the three experimental replicates. We took Fluc as a negative control to normalize the results in the three independent experiments. P1 significantly enhanced translation of the reporter protein (Fig. 9B). In contrast, the truncated version of P1 had no effect on translation (Fig. 9B). HC-Pro had an unexpected significant negative effect on translation that was even more intense when expressed as a P1/HC-Pro polyprotein (Fig. 9B). This result, remarkable by itself, showed that TEV HC-Pro possesses a translation inhibition activity. In retrospect, HC-Pro was not a good choice as a negative control in these experiments. Northern blot analysis of *in vitro* transcribed RNAs demonstrated that
differences in reporter protein accumulation were due to differences in translation efficiency because mRNA concentration was not significantly different in the reactions (Fig. 9E). These results suggest that TEV P1 and HC-Pro have some stimulatory and inhibitory effect on translation, respectively.

TEV translation depends on the IRES activity present in the viral 5’ UTR (49). To further investigate the effect of P1 and HC-Pro proteins on viral translation, we used a bicistronic reporter construct, as those usually used to analyze IRES activity (50-52). The bicistronic cassette consisted of the modified CPMV RNA-2 5’ UTR, the cDNA coding for Fluc, the TEV 5’ UTR, the cDNA coding for Venus, and the CPMV RNA-2 3’ UTR followed by a poly(A) tail (construct II, Fig. 9A and Fig. S4). Translation of the downstream cistron (Venus) depended on the IRES activity of TEV 5’ UTR. TEV P1 enhanced translation of upstream and downstream reporter proteins (Fig. 9C), indicating that the protein is able to stimulate translation driven by both UTRs, CPMV and TEV. In contrast, HC-Pro inhibited translation of both reporters, particularly the upstream but also the downstream. Finally, when both P1 and HC-Pro were present in the reaction, translation of the upstream reporter was inhibited, but interestingly the presence of P1 compensated the inhibitory effect of HC-Pro on translation of the downstream reporter. Taken together, these results suggest that TEV P1 enhances protein translation, particularly when it is driven from the viral 5’ UTR IRES, and that TEV HC-Pro inhibits translation.

DISCUSSION

In our work, we infected plants with a series of TEV recombinant clones in which the P1 protein was tagged either with a fluorescent protein to track P1 expression and localization during infection or with an affinity tag for specific purification in native conditions of protein complexes in which P1 is also a component. The proteomic analysis of these complexes fundamentally identified proteins from the ribosome 60S subunit (Table 1). But also, some proteins from the 40S subunit, such as ribosomal protein S6 previously shown to be strictly required for turnip mosaic potyvirus infection (53). Analysis of polysomes purified from infected tissues corroborated the physical association between TEV P1 and the plant cytoplasmic 80S ribosomes during infection (Fig. 8B), and more specifically with the 60S ribosomal subunits (Fig. 8C). In addition to proteins involved in translation, some other host proteins were also identified in association with TEV P1 in infected tissues (Table 1). It is worth noting heat shock protein 70 chaperones, previously shown associated to potyviral replication complexes (41, 45), and the glyceraldehyde-3-phosphate dehydrogenase.
(GAPDH) involved in replication of other positive-stand RNA viruses, like tomato bushy stunt tombusvirus (54).

Our experiments also demonstrate that TEV P1 is a nucleocytoplasmic protein that displays a dynamic intracellular localization during the infectious cycle (Fig. 3). At an early stage, the protein enters the nuclei and targets the nucleoli of infected cells. Later, P1 protein is exported from the nucleus to cytoplasm. These findings are supported by our identification of functional nucleolar localization and nuclear export signals in TEV P1 (Figs. 5 and 6). Moreover, our analysis of TEV mutants in the P1 cistron showed the in vivo relevance of these signals (Fig. 7). Inside the nucleolus, TEV P1 specifically localizes in the granular component, where final processing of pre-ribosomal particles takes place (55). In fact, TEV P1 nucleolar localization is similar to that of a genuine ribosomal protein (Fig. 4): A. thaliana RPL24B, a structural component of 60S ribosomal subunits. RNA viruses, including plant viruses, interact with the nucleolus to usurp host-cell functions, diverting nucleolar proteins to perform novel roles in the virus infectious cycle (56, 57). Potyviral NIa protein, more specifically the VPg domain, also contains nuclear and nucleolar localization signals. And nuclear and nucleolar localization of potato potyvirus A NIa was demonstrated to be essential for completion of the infectious cycle (58).

These findings suggest that potyviral P1 protein may play an unsuspected role in translation during the infectious cycle. TEV genomic RNA differs from a conventional host mRNA because it does not contain a 5'-cap structure. In addition, the TEV 5’ UTR has been shown to possess IRES activity (49, 59). The rate-limiting step in the initiation of translation of eukaryotic mRNA is the recognition of cap structure by eIF4E, the small subunit of eIF4F. In cap-independent translation of TEV, recruitment of the translation pre-initiation complex is not required because the viral IRES is able to directly bind the eIF4G and facilitates binding of the 40S subunit through interaction with other initiation factors (59-61). Sequences in the 3’ UTR of the pea enation mosaic tombusvirus have been shown to bind host ribosomal subunits assisting cap-independent translation of the viral mRNA (62). A model that may explain our experimental data is that, in the early stage of infection, potyviral P1 associates to the plant 60S ribosomal subunits to make them more competent in viral translation. This association may occur in the nucleolus during biogenesis of pre-ribosomal particles, but not necessarily because a TEV P1 mutation (Mt-IX) that abolishes nucleolar localization of the protein (Fig. 6B, IX) does not preclude binding to ribosomal 60S subunits during infection (Fig. 8D). P1 may mediate the recruitment of the 60S subunits to the viral translation initiation complex. CaMV P6 protein has also been shown to localize in the nucleolus and
interact with structural components of the 60S ribosome subunit (46, 63, 64). CaMV P6 is thought to trans-activate translation of the viral polycistronic pre-genomic RNA and its spliced versions. Recently, potyviral VPg has also been implicated in viral translation. Experiments based on transient expression of potato potyvirus A VPg in *N. benthamiana* plants have suggested that this viral protein stimulates translation of the viral mRNA and, at the same time, represses translation of the host mRNAs (11).

In support of the above model, *in vitro* expressed TEV P1 stimulated translation of reporter proteins (Fig. 9). These experiments led to the serendipitous observation that TEV HC-Pro inhibited translation (Fig. 9). This inhibition may result from the recently reported eIF4E and eIF(iso)4E binding activity of potyviral HC-Pro that may prevent the formation of functional cap-dependent translation complexes required for host protein production, resulting in selective inhibition of host cell protein synthesis (65). By incorporating the inhibitory effect of potyviral HC-Pro to our model, the coordinated action of P1 and HC-Pro during infection may stimulate IRES-dependent translation of the viral mRNA and suppress the cap-dependent translation of the host mRNAs. However, it should be taken into consideration that support to the model comes from *in vitro* translation experiments using RNAs with naked 5’ ends, which may not accurately reflect the situation occurring *in vivo* during virus infection.

When a potyvirus first enters a plant cell, after de-encapsidation, the viral genomic RNA needs to be efficiently translated to produce the proteins that will take part in the different steps of the infectious cycle. The virus may gain translation efficiency in a critical early stage of infection by the stimulating effect of viral P1. This is consistent with our observation that TEV P1 accumulates transitorily at the early stage of infection (Fig. 1). Since gene expression in potyviruses produces equimolar amounts of most proteins, this observation suggests a protein degradation mechanism to regulate P1 expression. The role we propose for potyviral P1 in our model is compatible with previous observations indicating that this cistron is dispensable for viral infection and that null P1 mutants can be rescued by expressing P1 protein in *trans*. Nonetheless, many viral proteins are multifunctional and P1 most probably plays multiple roles during infection. A recent work shows that the hypervariable amino-terminus of plum pox potyvirus P1 protein modulates replication and host defense response (66). Also, a transient suppression of host gene expression was reported exclusively at the infection front of pea seed-borne mosaic potyvirus (67). As P1 shows the same accumulation dynamics at the infection front, this protein may also be related with this process.

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We thank Verónica Aragonés and Teresa Cordero for excellent technical assistance. We thank C. Douglas Grubb (Leibniz-Institut für Pflanzenbiochemie, Germany) for critical review of the manuscript. Proteomic analysis was performed in the proteomics laboratory of Centro de Investigación Príncipe Felipe de Valencia, a member of Spanish ProteoRed. This work was supported by grant BIO2011-26741 from the Spanish Ministerio de Economía y Competitividad. F.M. was the recipient of a pre-doctoral fellowship from Universidad Politécnica de Valencia.

REFERENCES


Table 1. Refined list of identified host and viral proteins associated with TEV P1 protein in *N. benthamiana* infected tissue.

<table>
<thead>
<tr>
<th>Functional category</th>
<th>Protein hits&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Structural constituent of ribosome (60S)</td>
<td>Ribosomal proteins L3, L4, L6, L7, L10,</td>
</tr>
<tr>
<td></td>
<td>L11, L13, L18, L21, L23, L24, L27, L32,</td>
</tr>
<tr>
<td></td>
<td>L35 and P0</td>
</tr>
<tr>
<td>Structural constituent of ribosome (40S)</td>
<td>Ribosomal proteins S6 and S23</td>
</tr>
<tr>
<td>Translation</td>
<td>G-binding protein</td>
</tr>
<tr>
<td>Protein folding</td>
<td>HSP70-2, HSP70-3, CPHSC70-2 and BiP5</td>
</tr>
<tr>
<td>Structural constituent of cytoskeleton</td>
<td>Actin, tubulin alpha and beta</td>
</tr>
<tr>
<td>Photosystem</td>
<td>ATP synthase alpha and beta subunit</td>
</tr>
<tr>
<td>Metabolic process</td>
<td>GAPDH</td>
</tr>
<tr>
<td>Response to stress</td>
<td>Tudor-SN</td>
</tr>
<tr>
<td>Viral</td>
<td>TEV P1, HC-Pro and CP</td>
</tr>
</tbody>
</table>

<sup>a</sup>Score, Genbank accession numbers and full names of the proteins are in Table S1 (see supplemental materials).
LEGENDES TO THE FIGURES

FIG 1 Expression of P1 protein during TEV infectious cycle. (A) Schematic representation of recombinant TEV infectious clones: TEV-wt, TEV-VenusP1, TEV-GFP and TEV-VenusP1-mCherryNIb and TEV-TSTP1. Lines represent viral 5' and 3' UTRs, and white boxes represent viral cistrons P1, HC-Pro, P3, 6K1, CI, 6K2, VPg, NlaPro, Nlb and CP, as indicated. P3N-PIPO is represented by white (P3N) and dashed (PIPO) boxes. Light green, dark green and red boxes represent Venus, GFP and mCherry cDNAs, as indicated. The blue box represents Twin-Strep-tag (TST) cDNA. (B) Fluorescence images taken under a stereomicroscope of N. benthamiana leaves showing TEV-GFP and TEV-VenusP1 infection foci at 3 dpi. Scale bars correspond to 500 and 50 μm at low and high magnification, respectively. (C) Fluorescence images taken under a stereomicroscope of a TEV-VenusP1-mCherryNIb infection focus on an N. benthamiana leaf at 3 dpi. Scale bar corresponds to 50 μm. (D) Time-course analysis of TEV TSTP1 and CP accumulation in the third leaf of N. benthamiana plants above the leaf agroinoculated with TEV-TSTP1. Samples taken at different days post-inoculation (dpi), as indicated, were separated by SDS-PAGE and analyzed by western blot. (E) Plots of TEV TSTP1 and CP accumulation (measured by western blot analysis in arbitrary units, a.u.) versus dpi in the third leaf of N. benthamiana plants above the leaf agroinoculated with TEV-TSTP1. Bars indicate the standard deviation of the measures taken in triplicate plants.

FIG 2 N. benthamiana plants inoculated with various recombinant TEV clones. (A) Pictures taken at 8 dpi of representative N. benthamiana plants mock-inoculated and inoculated with TEV-wt, TEV-VenusP1, TEV-GFP, TEV-VenusP1-mCherryNIb and TEV-TSTP1, as indicated. (B) Analysis by RT-PCR of the viral progeny in plants infected by various recombinant TEV clones. Viral cDNAs were amplified by RT-PCR from RNA preparations from infected tissues at 10 dpi. Amplification products were analyzed by electrophoresis in 1% agarose gels, followed by ethidium bromide staining. Products of the upper and lower gels were amplified with primer pairs flanking the P1 (primer I, 5'-TTATTGCATGCCTAAGGATTCC-3'; and primer II, 5'-AGGAACGCCTCTCTATTAAATG-3') and the NIb (primer III, 5'-CTATTGCAGCAATTTAAATCATTTC-3'; and primer IV, 5'-CTTTGCGGAAGGGTGGAGCCCCG-3') cistrons, respectively. Lane 1, RT-PCR negative control; lanes 2 to 11, RT-PCR products from RNA preparations from tissues from...
individual plants infected by TEV-wt (lane 2), TEV-TSTP1 (lanes 3 to 5), TEV-VenusP1 (lanes 6 to 8) and TEV-VenusP1-mCherryNlb (lanes 9 to 11); lane 12, DNA marker ladder with the size in kbp of some of the components on the right.

FIG 3 P1 and Nlb subcellular localization during the TEV infectious cycle. (A) Schematic representation of an infection focus with cells (I to V) at different stages of infection. (B) Green and red fluorescence images taken under a confocal microscope of *N. benthamiana* leaf cells infected by TEV-VenusP1-mCherryNlb at 3 dpi. I to V indicate the position of the cells in the infection focus, from the periphery to the epicenter. Each series includes the green and red fluorescence images at low and high magnification and a merged image at high magnification. Scale bars correspond to 16 and 8 μm at low and high magnification, respectively.

FIG 4 Co-localization of *A. thaliana* AtFib2 and AtRPL24B nucleolar marker proteins and P1 during TEV infection. *N. benthamiana* leaves were inoculated with TEV-VenusP1 virions and infiltrated with *A. tumefaciens* cultures to transitorily express AtFib2mRFP and AtRPL24BmCherry. Green and red fluorescence images were taken under a confocal microscope three days later. A merged image is also shown. (A) Transient expression of AtFib2mRFP in a cell infected by TEV-VenusP1. Arrows point to nucleolus (No) and a Cajal body (CB). (B) Transient expression of AtRPL24B in a cell infected by TEV-VenusP1. (C) Schematic representation of a plant nucleolus for a better interpretation of results: nucleolar cavity (NC), fibrillar component (FC), dense fibrillar component (DFC) and granular component (GC). Scale bars correspond to 8 μm.

FIG 5 Identification of a nucleolar localization signal (NoLS) and a nuclear export signal (NES) in TEV P1 sequence (Genebank accession number ABJ16044). (A) NoLS prediction per residue displayed by NoD algorithm. (B) NES prediction in the 65 carboxy-terminal amino acids of TEV P1 displayed by NetNES algorithm. (C) Amino acid sequence of TEV P1 with the predicted NoLS and NES underlined and highlighted over pale and dark gray background, respectively.

FIG 6 Transient expression of Venus, VenusP1 and a series of VenusP1 mutants in *N. benthamiana*. Leaves were infiltrated with *A. tumefaciens* cultures to express the different constructs. (A) Schematic representation of the expressed constructs (I to X). The green and
white boxes represent Venus and TEV P1 cDNAs, respectively. The gray and black boxes represent TEV P1 NoLS and NES, as indicated. The black arrow and hexagon represent CaMV 35S promoter (P35S) and terminator (t35S), respectively. Black lines represent the *Cowpea mosaic virus* (CPMV) RNA-2 5’ and 3’ UTR, as indicated. Mutations in NoLS and NES sequences are represented in red. (B) Green fluorescence images of selected cells taken under a confocal microscope three days after infiltration with constructs I to X. Some cells were imaged at two different magnifications. Scale bars correspond to 20 and 8 μm at low and high magnification, respectively.

**FIG 7** Effect of mutations in the P1 cistron on TEV infectivity and accumulation. (A) Number of symptomatic *N. benthamiana* plants versus days post-agroinoculation of wild-type (WT) and P1 mutant TEV clones including the Ros1 marker. (B) Viral load at 3 days after symptoms emerged, measured as anthocyanin accumulation (absorbance at 530 nm), in plants infected by WT and P1 mutant TEV clones including the Ros1 marker. Error bars indicate the standard deviation of the three sampled plants. WT and mutant P1 (Mt-VII, Mt-VIII, Mt-IX, Mt-X, Mt-X+NlaPro, Mt-H214A and Mt-H214A+NlaPro) sequences are in supplemental material (Fig. S3).

**FIG 8** Identification of host proteins physically associated to P1 during TEV infection. TSTP1 was purified from *N. benthamiana* tissues infected with TEV-TSTP1 by liquid chromatography using a Strep-tactin column in native conditions. (A) Proteins eluting from the column were precipitated, separated by SDS-PAGE and the gel stained with Coomassie Blue. Lane 1, protein standards with their molecular masses in kDa on the left; lane 2, negative control of proteins purified from tissues infected by TEV-wt; lane 3, proteins purified from tissues infected by TEV-TSTP1. The lower panel corresponds to a western blot analysis using an anti-TST antibody. (B, C and D) Fractionation of polysomal preparations from *N. benthamiana* tissues infected by TEV-TSTP1 (B and C) or TEV-TSTP1 (mutant Mt IX) (D). A polysomal preparation from tissue infected with TEV-TSTP1 was non-treated (B) or treated with EDTA (C) and subjected to centrifugation in sucrose gradients. The fractionated gradient profiles were obtained by measuring the optical density of the different fractions at 254 nm. Ribosomal 18S and 28S RNAs were detected in the fractions of the gradients by agarose electrophoresis and staining with ethidium bromide. The presence of TSTP1 was revealed by SDS-PAGE separation and western blot analysis. The positions of the 80S ribosomes and the 60S and 40S ribosomal subunits are indicated in the profile. The
positions of the viral TSTP1 and the ribosomal 28S and 18S RNAs are indicated. (D) Two independent polysomal preparations from tissues infected with TEV-TSTP1 (Mt-IX) were treated with EDTA and fractionated by centrifugation in sucrose gradients. Ribosomal RNAs and TSTP1 (Mt-IX) were detected as indicated in (B) and (C). Lane 1 to 3, crude extracts from a non-inoculated plant and two independent plants infected with TEV-TSTP1 (Mt-IX), respectively; lanes 4 and 5, high speed sediments previous to fractionation from both infected tissues; lanes 6 and 8, and 7 and 9, peak fractions with the ribosomal 40S (lanes 6 and 8) and 60S (lanes 7 and 9) subunits for both infected samples, respectively. The positions of the viral TSTP1 (Mt-IX) and the ribosomal 28S and 18S RNAs are indicated.

FIG 9 Effect of TEV P1 on in vitro translation of reporter systems. (A) Schematic representation of the monocistronic and bicistronic reporter cassettes and the cassettes to express firefly luciferase (Fluc), a truncated form of TEV P1 (ΔP1; 155 initial amino acids), P1, HC-Pro and the P1/HC-Pro polyprotein. The black and gray rectangles represent the SP6 bacteriophage promoter and the poly(A) tail, respectively. Pale and dark gray boxes represent Fluc and HA-tagged Venus cDNAs, respectively. White boxes represent cDNAs corresponding to ΔP1 (non-translated part due to three stop codons inserted in frame is dashed), P1 and HC-Pro, as indicated. Black lines represent the CPMV RNA-2 5' and 3' UTR, as indicated. TEV 5' UTR is represented by a black rectangle, as indicated. (B) Production of Venus in in vitro translation reactions in which Fluc, TEV ΔP1, P1, HC-Pro and P1/HC-Pro were co-translationally produced. In vitro translation products in the presence of [35S]Met were separated by SDS-PAGE and quantified by phosphorimager analysis. Venus amounts were normalized by those obtained in the reaction co-translating Fluc. (C) Production of upstream Fluc (pale gray bars) and downstream Venus (dark gray bars) in in vitro translation reactions in which TEV ΔP1, P1, HC-Pro and P1/HC-Pro were co-translationally produced. The Fluc and Venus amounts were normalized by those obtained in the reaction co-translating TEV ΔP1. In (B) and (C) error bars indicate the standard deviation in three independent experiments and the different letters over the columns indicate a significant statistical difference (least significant different, LSD test p < 0.05). (D and E) Analysis of proteins and reporter mRNA produced during the coupled in vitro transcription-translation experiment. Two aliquots of the coupled in vitro transcription-translation reactions were taken at 40 min for protein and reporter mRNA analysis. (D) Proteins were separated by PAGE (12.5% polyacrylamide, 0.05% SDS), the gel dried and the proteins visualized by autoradiography. (E) RNAs were separated by electrophoresis in an agarose gel under
denaturing conditions, electroblotted to a positively charged nylon membrane and the Venus mRNA detected with a complementary RNA probe labeled with $^{32}$P. Lanes 1 to 5, reactions programed to produce Venus reporter and Fluc (lane 1), TEV P1 (lane 2), a truncated form of TEV P1 (ΔP1, lane 3), TEV HC-Pro (lane 4) or the P1/HC-Pro polyprotein (lane 5). The positions of the different proteins are indicated on the left and right of panel D. Note that the P1/HC-Pro polyprotein partially self-cleaves during the reaction. The position of the Venus reporter mRNA is indicated on the right of panel E.
NoLS predictions per residue

NetNES 1.1: predicted NES signals in sequence

>C>TEV P1 (Genbank no. ABJ16044)
MALIPTCTVMNLKEKYYFGARMACVTGC4MAAAGNC5LKEKREFTRRAIMKWKVPFGEDYI1TEDALDYPPFLC6LEDDDEEMNYLGRLKAMRAAMK4KVSNNKRLRKK45K1Y
GRRSIVKVERKVERKVDTDAAVIDE4A4TQTQVENSMFREKKQNNLFAATLSN
VYK7771IVRLEMMKWUI1DDDSVVARKVKRRFG2PVQFLFASVVRMM4GRRKVEQILIDN
M4QETTLDLAARFKNERVQGSKLTGSSKQVPPQGTYTPA4WTRK3GMFIVGR3DGMLV44AR4YTTFAVCNSMT5V