Transgenic Expression of Tobacco mosaic virus Capsid and Movement Proteins Modulate Plant Basal Defense and Biotic Stress Responses in Nicotiana tabacum

G. Conti,1,2 M. C. Rodriguez,1,2 C. A. Manacorda,1 and S. Asurmendi1,2

1 Instituto de Biotecnología, CICYVYa-INTA, Hurlingham, Buenos Aires, Argentina; 2 Consejo Nacional de Investigaciones Científicas y Técnicas (CONICET), Buenos Aires, Argentina

Submitted 31 March 2012. Accepted 14 June 2012.

Plant viruses cause metabolic and physiological changes associated with symptomatic disease phenotypes. Symptoms involve direct and indirect effects, which result in disruption of host physiology. We used transgenic tobacco expressing a variant of Tobacco mosaic virus (TMV) coat protein (CP142W) or movement protein (MP), and a hybrid line (MP×CP142W) that coexpresses both proteins, to study the plant response to individual viral proteins. Findings employing microarray analysis of MP×CP142W plants and silenced mp×cpT42W* controls revealed that altered transcripts were mostly downregulated, suggesting a persistent shut-off due to MP×CP142W expression. Next, we showed that MP triggered reactive oxygen species (ROS) accumulation, reduction of total ascorbate, and expression of ROS scavenging genes. These effects were enhanced when both proteins were coexpressed. MP and MP×CP142W plants showed increased levels of salicylic acid (SA) and SA-responsive gene expression. Furthermore, these effects were partially reproduced in Nicotiana benthamiana when GMP1 transcript was silenced. CP142W seems to be playing a negative role in the defense response by reducing the expression of PR-1 and RDR-1. MP and MP×CP142W transgenic expression promoted a recovery-like phenotype in TMV RNA infections and enhanced susceptibility to Pseudomonas syringae and Sclerotinia sclerotiorum. The individual effects of viral proteins may reflect the ability of a virus to balance its own virulence.

Viral diseases cause a great diversity of symptoms, producing irreversible yield losses in cultivated plants. Plant viruses often induce a number of common metabolic and physiological alterations that are probably the cause of the symptomatic phenotypes of disease. Virus-susceptible host interactions result in interference or competition for host resources and involve direct effects, which contribute to the establishment of a systemic infection, and side effects, which do not contribute to the success of the infection but indirectly disrupt host physiology. Plant gene expression is altered during virus infection and it has been reported that, like animal viruses, plant viruses are capable of reprogramming and disrupting host gene expression in a phenomenon known as “shut off” (Aranda and Maule 1998; Havelda et al. 2008). Genes involved in many cellular processes such as phytohormone signaling, cell cycle control, and endogenous transport of macromolecules and cell-wall- and plastid-related genes, among others, are either induced or repressed upon infection (Bazzini et al. 2011; Pallas and Garcia 2011; Shimizu et al. 2007; Whitham et al. 2006). Some of the distinctive gene expression changes resemble those observed in cellular stress and developmental defects (Whitham et al. 2006). The stress-like responses are characterized by the induction of heat-shock proteins (Aranda et al. 1996; Jockusch et al. 2001; Mayer 2005) and changes in the expression of several genes involved in reactive oxygen species (ROS) detoxifying mechanisms such as catalases, superoxide dismutases, peroxidases, and other redox-related genes (Li et al. 2005; Riedle-Bauer 2000).

The development of a systemic defense response triggered by a local ROS signal is mediated by salicylic acid (SA) production and by the induction of pathogenesis-related (PR) genes and other genes associated with plant disease defenses (Jones and Dangl 2006; Thimm et al. 2004; van Loon et al. 2006). The induction of many PR genes (Edreira 2005; van Loon et al. 2006; Yang et al. 1997) and other coregulated transcripts are involved in defense-like responses, which are usually mediated by defense hormones, predominantly SA (Loake and Grant 2000; Vlot et al. 2009). Although the induction of PR proteins is widely associated with disease defenses, none of them are known to have any antiviral effects (Cutt et al. 1989; Linhorst et al. 1989; Palukaitis et al. 2008).

Despite impressive advances in the study of plant resistance to pathogens, little is known about the plant response in a compatible virus interaction. In tobacco, Tobacco mosaic virus (TMV) replication is inhibited by SA treatment; which results in a reduced viral accumulation and a delayed appearance of disease symptoms (Naylor et al. 1998; Nie 2006). In compatible host–virus interactions, the expression of several defense-related genes is induced by an SA-dependent signaling pathway (Huang et al. 2005; Whitham et al. 2003). Moreover, the treatment of tobacco with SA restricts TMV movement in epidermal tobacco cells (Murphy and Carr 2002). There is a limited understanding of the global response taking place in the reaction of compatible hosts to viral infections owing to its complexity. For instance, the reaction depends on various features such as the demands of the virus, host defenses, host stress factors, cellular responses, and local or systemic responses. Transgenic host plants expressing viral proteins can be used as a simplified experimental system that has the potential to discriminate the effects of individual gene products from the impact of virus replication (Maule et al. 2002). For example, the ectopic expression of the Cauliflower mosaic virus P6 gene in

*The e-Xtra logo stands for “electronic extra” and indicates that three supplementary figures and two supplementary tables are published online. Also, Figures 1, 2, 4, 5, and 6 appear in color online.
its host species, Arabidopsis, has been shown to elicit symptom-like effects such as late flowering and mild chlorosis (Cecchini et al. 1997; Zijlstra et al. 1996). In a similar approach, the expression of TMV 126-kDa protein fused to green fluorescent protein (GFP) in N. tabacum produces an increased susceptibility to some viruses (Harries et al. 2008).

Transgenic tobacco plants expressing TMV coat protein (CP) fail to develop symptoms and to accumulate virions when inoculated with TMV (Abel et al. 1986) owing to a phenomenon called CP-mediated resistance (CP-MR) (Beachy 1999). When the CP amino acid residue 42 (Thr) is mutated to Trp (CP<sup>T42W</sup>), there is an increase of CP 20S aggregate amounts that could be responsible for the enhanced effect on the host and on the CP-MR level (Asurmendi et al. 2007; Bazzini et al. 2007; Bendahmane et al. 1997). In addition, CP<sup>T42W</sup> restricts cell-to-cell spread by reducing the production of movement protein (MP) during TMV replication (Bendahmane et al. 2002), and it induces mild heterologous resistance to other viruses (Bazzini et al. 2006). Thus, several mechanisms of action must be taking place during the CP-MR phenomenon triggered by CP<sup>T42W</sup>. Inoculation with TMV RNA partially overcomes protection against virus infection in CP transgenic plants (Nelson et al. 1987); in contrast, CP<sup>T42W</sup> transgenic lines still show some degree of resistance. Constitutive expression of MP in tobacco plants showed that this 30-kDa protein interacts with the plasmodesmata, modifies their exclusion limit, and facilitates cell-to-cell spread of the virus during infection (Atkins et al. 1991; Deom et al. 1987, 1990; Wolf et al. 1989).

The TMV MP has been demonstrated to cause severe physiological alterations, such as carbohydrate partitioning and photosynthesis on host plants, when expressed under constitutive or tissue-specific promoters (Almon et al. 1997; Balachandran et al. 1997). Moreover, it is also very interesting to consider the outcome of the co-accumulation of CP and MP to achieve a better understanding of the infective process. For instance, in a previous study, the coexpression of MP and CP<sup>T42W</sup> on hybrid plants was accompanied by severe morphological changes, altered miRNAs accumulation, and poor fertility. In addition, it is worthy to note that no detectable silencing suppression activity in relation to defense signaling was detected on the mentioned proteins (Bazzini et al. 2007).

In this work, we applied a transgenic approach to study the disruption of the natural host physiology and the triggering of a defense response by individual TMV protein expression. We made use of transgenic Nicotiana tabacum lines expressing either a variant of TMV CP, a TMV MP, or both proteins. The coexpressing line was obtained by means of a genetic cross between the two single protein transgenic lines (this new line was called MP×CP<sup>T42W</sup>). To study the differential expression patterns resulting from the transgenic coexpression of MP and CP<sup>T42W</sup> in a TMV natural host, we performed a microarray analysis. We further focused the analysis on several array-based selected genes related to stress and defense pathways.

Next, we evaluated the accumulation level of ROS and defense hormones, such as SA and jasmonic acid (JA). Finally, we evaluated the impact on the defense system of these transgenic plants when challenged against a wide spectrum of pathogens, and we also performed functional assays to examine the influence of an individual gene in the phenotype and defense response observed in MP×CP<sup>T42W</sup> plants.

**RESULTS**

**The co-accumulation of TMV MP and CP<sup>T42W</sup> mediates global changes in gene expression.**

To characterize the gene expression changes caused by the coexpression of TMV capsid proteins and MP in N. tabacum ‘Xanthi’, we monitored global gene expression patterns with a Tomato 10K Affymetrix Microarray GeneChip comparing the isogenic transgenic lines MP×CP<sup>T42W</sup> (transgenic tobacco SX coexpressing MP and CP<sup>T42W</sup> genes) and mp×cp<sup>T42W</sup>*. The latter line was used as a control genotype, considering that local transgene insertions and genetic background are conserved between both lines but that the expression of the transgenes is silenced in mp×cp<sup>T42W</sup>* (Bazzini et al. 2007). The array contains more than 10,000 *Solanum lycopersicum* probe sets to monitor gene expression of more than 9,200 *S. lycopersicum* genes. It has already been verified that this array is useful for several solanaceous species, including tobacco, owing to the high degree of sequence conservation between transcripts (Rensink et al. 2005). The gene expression matrix was analyzed in two concatenated phases. First, we considered transcripts of the genes relevant to treatment differentiation, defined by P values < 0.10. In the second phase, we applied a P value correction to control false discovery rates (FDR) (P value < 0.05) to the set generated in the first instance, as previously reported by Benjamini and Hochberg (1995). Raw data of the experiment were deposited at Gene Expression Omnibus (GSE37905). The analysis yielded 28 genes showing statistically significant enhanced or reduced accumulation levels (Fig. 1A). The selected transcripts were categorized based on shared structural elements or inferred function making a comparison between tomato and tobacco genomes using BLAST (Table 1). The results comprised various subgroups, including genes related to oxidative stress and hormone response, signal transduction, transcriptional regulation, carbohydrate metabolism, and RNA
processing (Fig. 1B). These results were validated by reverse-transcriptase quantitative polymerase chain reaction (RT-qPCR) between the transgenic tobacco lines, confirming 16 of 23 genes tested (Supplementary Fig. S1); 5 genes were not tested because their PCR amplification failed using the available sequence data. Interestingly, most of the differentially expressed transcripts that were included in oxidative stress, hormone response, signal transduction, and metabolism functional categories were downregulated in plant lines coexpressing MP and CPT42W (Table 1). For the subsequent work, we focused on a group of genes related to oxidative stress and a second group associated with defense hormones. To deepen into the study of redox state and defense hormone regulation, we extended the analysis to a wider spectrum of genes and also to the parental transgenic CPT42W and MP lines and to wild-type SX tobacco plants.

Viral proteins increase ROS accumulation and modulate ascorbic acid contents and redox scavenging gene expression.

Considering that the MP×CPT42W line showed a reduced expression of genes involved in redox balance, we examined whether the transgenic expression of individual CPT42W and MP or the coexpression of both viral proteins altered the accumulation of ROS and ascorbic/dehydroascorbic acid levels (ASC/DHA) in leaf tissues. To achieve that aim, we performed histochemical staining to determine the accumulation of hydrogen peroxide (H₂O₂) and superoxide ions (O₂⁻) using diaminobenzidine (DAB) and nitroblue tetrazolium (NBT), respectively. Results showed that MP and MP×CPT42W transgenic lines accumulated elevated levels of H₂O₂ and O₂⁻ compared with wild-type plants. These effects were more pronounced in the coexpressing line MP×CPT42W, which also showed higher cell death (detected by trypan blue staining) (Fig. 2A). Finally, the ASC/DHA measurements demonstrated that MP×CPT42W accumulated reduced levels of both antioxidant compounds (Fig. 2B). Altogether, these results suggest a synergic effect between these viral proteins enhancing an stress-like response phenotype in MP×CPT42W compared with MP plants. As expected, silenced mp×cpT42W* plants showed normal ROS and ASC/DHA levels, confirming that the effect is exclusively due to viral protein expression. On the other hand, CPT42W transgenic tobaccos behaved like wild-type controls, suggesting that CP alone fails to produce a stress-like phenotype similar to that observed with individual expression of MP or the combined expression of both transgenic proteins.

Based on the results of Figures 1 and 2A and B, we also performed RT-qPCR in CPT42W, MP, wild-type controls (SX), and systemic tissue of TMV-infected tobacco plants. The aim was to broaden the analysis made with MP×CPT42W, to include a larger set of redox-related genes. The analysis of TMV-infected tobacco plants was performed early in infection (at 6 days postinfection [dpi], with no systemic viral accumulation) and in late infection (at 18 dpi, with high virus accumulation in systemic tissue) (Supplementary Fig. S2).

Results showed that GDP-mannose pyrophosphorylase 1 (GMP1) (SGN-U442525) transcript levels were reduced in MP×CPT42W and upregulated in TMV-infected plants at 18 dpi (Fig. 2C). Detoxifying cytosolic ascorbate peroxidase (cAPX1) (SGN-U448855) did not change its expression level in any of the transgenic lines or TMV-infected plants at 6 dpi but it was upregulated in TMV-infected plants at 18 dpi (Fig. 2C). Chloroplast superoxide dismutase 2 (CSD2) (SGN-U313819) transcript levels were reduced in MP×CPT42W but it was upregulated in TMV infections at 18 dpi. Expression levels from all the transcripts measured were not altered in CPT42W and silenced mp×cpT42W* (Fig. 2C). Finally, the relative level of detoxifying

<table>
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<td>↑</td>
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^a Arrows ↓ and ↑ indicate downregulated or upregulated transcripts, respectively.
alternative oxidase (AOX) (SGN-U450832) transcript was lower in MP×CP^{T42W} and was not altered by TMV infection (Fig. 2C). The reduced levels of GMP1, CSD2, and AOX could play an important role in the enhanced stress-like phenotype observed in MP×CP^{T42W}.

In accordance with previously reported works (Fodor et al. 1997; Li et al. 2005; Wise et al. 2007), in this study, redox-related genes were induced at late stages of TMV infections (18 dpi) (Fig. 2C). However, even though an increment of detoxifying gene expression was expected, the sole MP expression produced no increase of the transcription levels of such genes, suggesting that this protein somehow deregulates the ROS scavenging system in host plants. Interestingly, the coexpression of MP and CP^{T42W} showed a stronger phenotype associated with ROS increments, further reducing the detoxifying enzyme levels; which, in turn, enhanced this particular phenotype.

MP expression increases SA levels and induces SA-responsive genes, whereas CP^{T42W} expression produces a partial antagonistic effect.

As stated in the differential transcript profiling of MP×CP^{T42W} tobacco plants, the genes related to hormonal response (SA, ethylene, and auxins) exhibit altered expression patterns. It has already been reported that SA plays a crucial role in plant disease and pathogen resistance (Durrant and Dong 2004; Vlot et al. 2009). In particular, SA levels are increased in TMV-resistant tobacco, which is paralleled with a raise in the expression of PR genes (Grant and Lamb 2006; Malamy et al. 1990; Vlot et al. 2009). For these reasons, in the following experiments, we focused on the study of SA levels and the expression of SA-responsive genes. First, we quantified SA and JA levels in the transgenic lines. Whereas SA was enhanced, JA was markedly reduced in leaves of MP and coex-
pressing MP×CP T42W plants (Fig. 3A). As a second strategy, we performed RT-qPCR to evaluate the expression levels of SA-responsive genes PR-1 (SGN-U444943), PR-2 (SGN-U439941), and PR-5 (SGN-U446829). In agreement with the SA level observed (Fig. 3A), all of these genes were upregulated in MP and MP×CP T42W plants (Fig. 3B), and no increase was detected in the silenced control mp×cp T42W* line. Unexpectedly, a reduction in PR-1 transcripts was found in CP T42W

Fig. 3. Characterization of defense-like phenotypes. A, Salicylic acid (SA) and jasmonic acid (JA) levels determined by high-performance liquid chromatography in transgenic lines of a variant of Tobacco mosaic virus (TMV) coat protein (CP T42W), movement protein (MP), and a hybrid line coexpressing both (MP×CP T42W) compared with wild-type control plants (SX). B, Relative expression levels of SA-responsive genes PR-1, PR-2, and PR-5 and genes responsible for silencing defense against viral RNAs, RDR1, and RDR6 in transgenic tobaccos and wild-type SX controls, CP T42W, MP, MP×CP T42W, the silenced control (mp×cp T42W*) and TMV-infected plants at 6 and 18 days postinfection (dpi), determined by reverse-transcriptase quantitative polymerase chain reaction. The mean relative transcript level ± standard error of five replicates is shown. Asterisks indicate significant differences between transgenic plants, compared with the wild-type SX (* and ** indicate P values < 0.05 and 0.01, respectively).
plants. The induction of most of the SA-responsive tested transcripts was also observed in TMV-infected plants at 18 dpi. 

RDR1 (referred to as NtRdRP1) (SGN-U431860), a SA-responsive gene directly involved in RNAi defense against viruses (Xie et al. 2001), was upregulated in TMV-infected plants at 18 dpi but, surprisingly, it was reduced in CP^{T42W} and MP×CP^{T42W} lines. Additionally, we found that transgenic viral protein expression did not alter RDR-6 (SGN-U437134) ex-

Fig. 4. Effects of Tobacco mosaic virus (TMV) RNA infection. A, Bar graphs show the percentage of TMV-infected plants (enzyme-linked immunosorbent assay [ELISA] positive) during the infection time course. Line graphs show the accumulation level of coat protein (CP) (mean values ± standard error of eight replicates per plant line) quantified by ELISA for samples collected at 0, 6, 8, 10, 15, 20, and 25 days postinfection (dpi). Gray shadow boxes highlight the time point were statistical analysis is shown, comparing viral accumulation levels between transgenic plant lines and SX controls. Asterisks indicate significant differences between transgenic plants compared with the wild-type SX (*, **, and *** indicate P values < 0.05, 0.01, and 0.005, respectively). Data shown are from a representative assay of three independent replicates. B, Differential symptoms displayed by all the transgenic plant lines and wild-type control plants (SX) at 10 dpi as indicated in the figure and recovery-like phenotype observed in 25 dpi movement protein (MP) tobacco (bottom panel). The black arrow on the left indicates a leaf with severe TMV-symptoms and the right arrow shows a leaf with mild symptoms demonstrating the recovery phenotype. Similar results were obtained in three independent assays.
expression, another antiviral gene implicated in post-transcription gene silencing (Curaba and Chen 2008), but TMV systemic infection did induce its expression.

Altogether, these results suggest that the MP behaves as a defense elicitor inducing a defense signal mediated by SA. In agreement with the stress-like response observed in Figure 2, it seems that the constitutive expression of MP triggers defense-like responses. A similar but stronger effect was demonstrated in MP×CP T42W coexpressing plants. However, the diminishing effect of CP T42W on the PR-1 and RDR-1 expression levels suggests a role for this viral protein as a negative modulator of specific defense-like responses.

Transgenic expression of MP promotes a recovery-like phenotype in infections with TMV RNA.

The changes of the stress and defense responses produced in the transgenic lines could alter the course of a pathogen infection. To test the effects of TMV CP and MP coexpression on the infectivity of plant viruses, each transgenic line and the wild-type control SX were inoculated with TMV RNA that is not affected by CP-MR. The virus accumulation in the systemic tissue was verified by enzyme-linked immunosorbent assay (ELISA) at different time points upon inoculation (6, 8, 10, 15, 20, and 25 dpi) and the progress and severity of symptoms were also monitored. (Fig. 4A and B.). All wild-type SX plants showed virus accumulation at 10 dpi and displayed the highest virus accumulation level at 25 dpi. As expected, the CP T42W transgenic line showed an initial mild resistance to TMV RNA (Bazzini et al. 2007), showing statistically significant lower accumulation at 10 dpi. However, at 25 dpi, 80% of the inoculated plants became infected, reaching a moderate viral accumulation and milder symptoms. In turn, the mp×cp T42W line was highly resistant: only 60% of the plants showed detectable virus at late stages, presenting statistically low virus accumulation and almost no symptoms at 25 dpi. Interestingly, despite the expression of CP T42W, MP×CP T42W plants were susceptible to TMV RNA infection, and 100% of the plants were infected and showed viral titles similar to SX controls until approximately 10 dpi. At late stages (25 dpi), there was an infection decrease in virus accumulation and fewer plants showed detectable levels of virus, demonstrating a partial recovery on systemic viral infection accompanied by a reduction in disease symptoms (Fig. 4A, gray shadow boxes). A similar effect was observed in the MP plant line, which showed even higher susceptibility than SX controls at early stages (until 10 dpi), reaching a total of 80% of the plants with detectable virus at 6 dpi and the highest virus accumulation at 10 dpi. At late stages (25 dpi), MP-infected plants showed a strong recovery process, as indicated by a reduction from 100 to 60% of the plants with detectable virus accompanied by significantly lower virus accumulation (Fig. 4A, gray shadow boxes) in the infected tissue. In summary, MP×CP T42W and MP plant lines are somehow more susceptible to TMV RNA at early stages of infection and more resistant in late stages because of partial recovery and symptom-shifting.

MP expression and coexpression of MP and CP T42W enhance susceptibility to hemibiotrophic and necrotrophic pathogens.

Inoculations with hemibiotrophic Pseudomonas syringae pv. tabaci and necrotrophic fungus Sclerotinia sclerotiorum were performed as an attempt to examine the effect of TMV CP and MP on the infectivity of pathogens with diverse lifestyles and with differences on the defense pathway induction. Leaves infiltrated with P. syringae pv. tabaci developed necrotic lesions in all plant lines and wild-type controls; however, the lesions observed in MP and MP×CP T42W plants were more severe (Fig. 5A). After 48 h, leaf discs from infected tissue were collected and incubated in King’s B medium. Plate count showed higher bacterial growth rates in MP and MP×CP T42W transgenic plants (Fig. 5B). Differences in the development of disease symptoms caused by S. sclerotiorum were also observed (Fig. 5C). The average necrotic lesion size in infected leaves of control and transgenic lines was measured. In agreement with results observed with P. syringae infection, MP×CP T42W transgenic lines showed statistically significant wider necrotic lesions 6 days after S. sclerotiorum inoculation and MP lines showed a similar but not significant tendency (Fig. 5D). Taken together, these results showed that MP and MP×CP T42W transgenic lines had an enhanced susceptibility to P. syringae pv. tabaci and S. sclerotiorum.
The silencing of GMP1 produced stress-like responses similar to those from the MP×CP T42W phenotype and enhanced resistance against TMV infections in N. benthamiana.

According to data from the expression profile and RT-qPCR ratios, GMP1 was downregulated in MP×CP T42W plants. Ascorbic acid contents were significantly reduced in these plants (Fig. 2B). As previously reported, the Arabidopsis vtc-1 mutant (knock-down mutant of GMP1) is deficient in ascorbic acid content, has a retarded growth and flowering (Conklin et al. 2000; Pastori et al. 2003), accumulates high ROS, exhibits increased cell death rates and elevated SA levels, induces PR-1 and PR-5 expression, and has an enhanced basal resistance against P. syringae infection (Colville and Smirnoff 2008; Mukherjee et al. 2010; Pavet et al. 2005). Taking all these findings into account, we explored in detail a possible involvement of GMP1 in the stress and defense responses observed in MP×CP T42W plants. To analyze this putative effect, a Tobacco rattle virus (TRV)-based virus-induced gene silencing (VIGS) system was employed for mimicking GMP1 downregulation in transgenic MP×CP T42W plants. pTRV-gmp1, carrying a partial N. tabacum GMP1 sequence (506 bp), and pTRV-gfp controls were infiltrated in N. benthamiana plants and systemic GMP1 silencing was confirmed by RT-qPCR (Fig. 6A). The gmp1-silenced plants dramatically reduced growth and showed higher ROS levels, detected in the leaf tissue by staining with DAB and NBT (Fig. 6B and C). These results confirmed that GMP1 plays an important role in the ROS scavenging system of N. benthamiana and suggested that GMP1 could be part of the stress-like phenotype observed in MP×CP T42W plants. When gmp1-silenced N. benthamiana plants were infected with TMV, an enhanced resistance was observed at late stages of infection (15 dpi) (Fig. 6D and E). This effect could be somehow parallel to that observed in MP and MP×CP T42W plant lines when infected with TMV RNA. Additionally, we observed similar results in the accumulation level of defense and stress response genes given by an increased accumulation of PR-1 and decreased levels of CSD2 (Fig. 6F). Increased tolerance against a low inoculum of the tobravirus Oiled-raped mosaic virus (ORMV) was also displayed in 6-week-old Arabidopsis vtc-1-mutants (Figs. 6G and H). Therefore, gmp1 silencing in both N. benthamiana and Arabidopsis vtc-1 mutants displayed an enhanced resistance to viral infections, which indicates that the earlier triggering of the defense response mediated by ROS produces an increased resistance in the plants.

DISCUSSION

A rapid viral replication in plant tissues involves the synthesis of large amounts of virus nucleic acids and proteins that must redirect host resources from normal cellular processes toward this viral replication process. It has already been reported that host demands can be modified by the virus through the specific downregulation of cellular gene expression (Aranda and Maule 1998). The effect, called host gene shut off, compromises some aspects of plant physiology and broad-spectrum defense response (Bazzini et al. 2011; Havelda and Maule 2000; Maule et al. 2002; Pallas and Garcia 2011). Shut off has been described as a transient effect developed within the virus replication cycle. However, Havelda and associates (2008) have demonstrated that viruses from some RNA virus families induce host gene transcript downregulation not directly in association with active viral replication. Furthermore, they have observed that the phenomenon persists for several weeks. In this study, we demonstrated that the stable transgenic coexpression of TMV CP T42W and MP could induce a gene-specific and persistent shut-off phenomenon. This effect is independent from viral replication and the depletion of host resources that occur during a systemic TMV infection, as evidenced by the insignificant accumulation of transgenic viral nucleic acids and proteins when compared with the protein level reached by the infection itself (Bazzini et al. 2007).

As clearly demonstrated by transcript profiling (microarray analysis and RT-qPCR), some genes required for ROS scavenging and defense signaling were downregulated in MP×CP T42W. It has been well documented that ROS are important in plant adaptation to both biotic and abiotic stresses (Apel and Hirt 2004; Mittler 2002). Under these conditions, ROS may play different roles. For example, they could be exacerbating damage or activating defense response signaling (Dat et al. 2000; Mittler et al. 2011). Consequently, the concentration of ROS has to be tightly regulated to maintain cell homeostasis. During compatible Cucumis sativus–Cucumber mosaic virus and Cucurbita pepo–Zucchini yellow mosaic virus interactions, plants displayed enhanced antioxidant capacities and inhibited local cell death by inducing superoxide dismutases, catalases, and ascorbate peroxidases, among others (Riedle-Bauer 2000). However, total ascorbate content and activities of scavenging enzymes were slightly and transiently reduced in TMV-inoculated leaves of N. tabacum before hypersensitive response (Fodor et al. 1997; Mittler et al. 1998; Yi et al. 1999). Thus, the suppression of the antioxidant system is important for local PCD induction in hypersensitive responses developed after localized pathogen recognition (Dat et al. 2003; Lorrain et al. 2003; Mittler et al. 1999; Yi et al. 2003). Immediately, compensatory mechanisms must necessarily be activated to stop PCD signaling and to maintain ROS formation (Bajda et al. 2009; Liao et al. 2012). These data are in agreement with our results, which showed that elevated transcript levels of cAPX1, CSD2, and GMP1 were produced in TMV-infected tobaccos at 18 dpi. Interestingly, we observed generalized and systemic decreases in total ascorbic acid content and reductions of GMP1, CSD2, and AOX transcript levels as a consequence of CP T42W and MP coexpression. In addition, the transcript levels of cAPX1 were unchanged in MP and MP×CP T42W plant lines. Altogether, these effects resulted in an enhanced stress response phenotype given by a high accumulation of ROS. Compensatory mechanisms to suppress toxic ROS levels were not activated, resulting in a strong ROS accumulation that could lead to the production of PCD micro-focuses in the MP×CP T42W transgenic line. Moreover, this line showed reduced growth and defective development (Bazzini et al. 2007). The downregulation of the transcription of scavenging enzyme genes could be mediated by the enhancement of MP function on the complex formed by MP and CP T42W (Asurmendi et al. 2004; Bazzini et al. 2007). A similar effect has been observed in pea plants inoculated with Plum pox virus (PPV) (Diaz-Vivancos et al. 2008). At 3 dpi, no changes were detected in the oxidative stress parameters, whereas decreases in chloroplast enzymatic mechanisms (ascorbate peroxidase and peroxidase) resulted in increased levels of H2O2 in this compartment. The stress response was higher at 15 dpi, as judged from exacerbated stress parameters and imbalances detected in the antioxidant system. However, the modulation of the expression of detoxifying and antioxidant enzymes has been shown to enhance basal defense against pathogens (Barth et al. 2004; Mittler et al. 1999; Pavet et al. 2005) and to induce the expression of plant defense proteins (Pastori et al. 2003; Pnueli et al. 2003).

There are additional regulatory functions for ROS in defense that may occur in conjunction with other signaling molecules, particularly with SA (Alvarez 2000; Durrant and Dong 2004). It has already been demonstrated that, in resistance-gene-mediated TMV-resistant tobacco plants, SA levels increase more than 20-fold in the inoculated leaves and approximately fivefold
Fig. 6. Effects of the GDP-mannose pyrophosphorylase 1 (GMP1) silencing on development, stress, and defense responses in *Nicotiana benthamiana*. **A**, Scheme of pTRV1 and pTRV2-gmp1 constructs used for *Tobacco rattle virus* (TRV) infection by agroinfiltration and reverse-transcriptase quantitative polymerase chain reaction (RT-qPCR) verification of GMP1 silencing 2 weeks after virus-induced gene silencing (VIGS). **B**, Effects of GMP1 silencing on plant growth and development 2 weeks after inoculation. **C**, Diaminobenzidine and nitroblue tetrazolium staining of GMP1-silenced leaves versus TRV-green fluorescent protein (gfp) control leaves. **D**, Differential symptom development of TRV-gfp control (left) and TRV-gmp1-silenced (right) plants inoculated with TMV at 10 days postinfection (dpi). **E**, Bar graphs showing relative levels of TMV coat protein (CP) accumulation in individual TRV-gfp control and TRV-gmp1 plants as determined by enzyme-linked immunosorbent assay (ELISA) test. **F**, Effects of GMP1 silencing on relative expression levels of salicylic-acid-responsive genes pathogenesis-related (PR)-1, PR-2, and PR-5 and detoxifying chloroplast superoxide dismutase 2 (CSD2) in TRV-gmp1 and TRV-gfp plants determined by RT-qPCR 2 weeks after inoculation. The mean relative transcript level ± standard error of five replicates is shown. Asterisks indicate significant differences (* and ** indicate $P$ values < 0.05 and 0.01, respectively). **G**, Mean relative accumulation levels of *Oiled-rape mosaic virus* (ORMV) CP (as determined by ELISA test) in 3- and 6-week-old *vtc1*-Arabidopsis mutants and Col0 controls at 7 dpi. Mean values were obtained from 10 plants per treatment. **H**, Differential symptom severity in 6-week-old *vtc1*-Arabidopsis mutants and Col0 control plants inoculated with ORMV at 7 dpi. Statistical differences are shown by *, **, and ***, indicating $P$ values < 0.05, 0.01, and 0.005, respectively.
in the systemic leaves. In addition, PR gene expression is also induced in these plants (Malamy et al. 1990). Interestingly, treatment of susceptible tobacco with SA causes a profound reduction in TMV accumulation. This effect occurs with no macroscopically visible cell death, and associated with ROS mediated signaling (Chivasa et al. 1997; Urbanek Krajnc et al. 2011). Enhanced levels of SA and decreased levels of JA were observed in all transgenic plant lines that overexpress MP, which suggests that at least this viral protein is sufficient to modulate the stress and defense responses in tobacco plants. In addition, it has been proposed that PR-1 proteins are not involved in transgenic TMV-CP-mediated resistance mechanism (Carr et al. 1989). Our findings demonstrated that a mutated version of CP (CP[T42W]) downregulates PR-1 and RDR-1 in transgenic plants. This effect is evident in MP×CP T42W for RDR-1 expression but not for PR-1. Therefore, CP[T42W] could be acting as a negative modulator of specific host defenses. This response is different from that elicited by MP expression; however, both responses could be derived from similar shut-off mechanisms with differential gene specificity for each viral protein.

The negative effect on defense activation described for CP[T42W] is in agreement with the reported abilities of CP to suppress or evade some plant responses by interacting with host factors (Callaway et al. 2001). Moreover, TMV CP facilitates TMV systemic movement by specifically interacting with a host component in N. tabacum (Hilf and Dawson 1993). Therefore, a tempting interpretation of our results point toward a role for the CP-mediated downregulation phenomenon of defense genes in the systemic viral movement.

The initial response triggered by TMV RNA infection in MP and MP×CP[T42W] lines is characterized by a rapid and systemic spread of the virus and also by an accelerated progression of symptoms. This first stage of the infection is followed by a recovery stage with a conspicuous reduction of both TMV accumulation and symptoms in systemic tissue. In addition, MP×CP[T42W] plants showed slightly reduced virus accumulation, which probably demonstrates a balance between rapid virus spread owing to a direct MP activity and a delay in virus systemic accumulation given by CP[T42W] expression. By contrast, control plants exhibited a sustained increase in TMV accumulation and symptom development. We hypothesize that the first stage (in which virus accumulates faster than in nontransgenic plants) could be produced by an enhanced spread of TMV transcripts caused by the expression of MP which, in turn, facilitates the spread of virus, as has been previously reported (Guenoune-Gelbart et al. 2008; Niehl and Heinlein 2011). It could also be argued that the enhanced expression of β 1-3 glucanases (PR-2) in MP-expressing lines might facilitate virus movement by degrading callose deposition in plasmodesmata (Baebler et al. 2011; Beffa et al. 1996; Epel 2009). In the second stage (in which virus accumulation decays and recovery of disease symptoms occurs), we hypothesize that the expression of MP promotes the control of the disease not only by means of enhancing the transport of the silencing signal, as has been previously proposed (Vogler et al. 2008), but also by inducing the activation of defense responses mediated by ROS and SA. Thus, the activation of basal defense responses and the enhancement of RNA-silencing given by MP transgenic expression could produce a combined effect that results in a reduced systemic viral spread and accumulation, giving rise to a partial recovery from infection. RNA-silencing and SA-dependent responses are suggested to play key roles in limiting Tomato ring spot virus spread in tobacco (Jovel et al. 2011). SA has also been proposed to enhance RNA-silencing antiviral defense against PPV in tobacco; accordingly, suppressors of gene silencing such as P1/HC-Pro would interfere with SA-mediated defense (Alamillo et al. 2006). The expression of RDR1 is induced by SA and in response to infection by many viruses, and it has been implicated in antiviral immunity by allowing the production of secondary small-interfering RNAs (Garcia-Ruiz et al. 2010; Xie et al. 2001). Despite all the evidence exposed that shows a connection between induction of RDR-1 and defense response mediated by SA, other mechanisms could be involved, as demonstrated in this work. For instance, our findings showed that the MP transgenic line did not induce RDR-1 expression and the MP×CP[T42W] line displayed a reduced RDR-1 transcript accumulation.

Plants trigger distinct defense responses depending on the lifestyle of the pathogen encountered. SA mainly induces defense against biotrophic pathogens that reproduce in live host cells, whereas JA acts mainly against necrotrophic pathogens that require killing host cells to grow and reproduce. Cross-talk between these defense pathways has been demonstrated to be essential to optimize the response against pathogens. Here, we showed that the expression of MP and the coexpression of MP and CP[T42W] result in enhanced susceptibility against P. syringae pv. tabaci and S. sclerotiorum infections. P. syringae pv. tabaci is a hemibiotrophic pathogen, with an initial biotrophic phase during which the infection is established, followed by a necrotrophic phase during which the pathogen completes its life cycle. Necrotrophic phases require production of large quantities of ROS and H₂O₂ for successful infections (Able 2003). In addition, necrotrophic pathogens such as S. sclerotiorum or Botrytis cinerea trigger hypersensitive cell death to facilitate Arabidopsis colonization (Govrin and Levine 2000). The enhanced susceptibility shown by MP and MP×CP[T42W] transgenic lines could be produced because both pathogens encountered an environment of high ROS accumulation that enabled them to initiate a necrotrophic phase. Another possibility could be that this accentuated susceptibility would be mediated by the activation of the SA-dependent response that is probably antagonistic to the specific response required for those particular pathogens. Similarly, Faize and associates (2011) have demonstrated that the overexpression of cytosolic Cu/Zn-superoxide dismutase and ascorbate peroxidase increase tolerance against P. syringae pv. tabaci and Agrobacterium tumefaciens in tobacco plants.

Finally, to gain a better insight into the mechanisms underlying the stress and defense outcomes in MP and MP×CP[T42W] transgenic lines, we analyzed the role played by GMP1 using a functional approach. This gene was not induced, although there were an ROS level increase in the MP plant line and it was reduced in the MP×CP[T42W] line. A similar phenotype, with high ROS accumulation, has already been reported in Arabidopsis vtc1 mutants (Foyer et al. 2007; Mukherjee et al. 2010; Pastori et al. 2003) and in gmp1-silenced N. benthamiana. Reduced levels of GMP1 have been reported to be sufficient to enhance basal defense mediated by SA and PR protein expression in Arabidopsis thaliana (Barth et al. 2004; Pau et al. 2005). In this study, an increased accumulation of PR proteins and an enhanced basal defense response against TMV were confirmed for gmp1-silenced N. benthamiana. A similar effect was observed in Arabidopsis vtc1 challenged against ORMV. The defense outcome evidenced in gmp1-silenced N. benthamiana, Arabidopsis vtc1, MP×CP[T42W], and MP suggests that plants are primed for defense against Tobamovirus spp. in a GMP1-dependent manner (directly or indirectly). It is important to mention that the recovery effect (late phenotype) showed in the MP×CP[T42W] and MP lines is the one considered here related to ROS response. However, the interaction is more complex, as judged by the differences observed in defense responses triggered against P. syringae. Therefore, according to the results obtained, different defense mechanisms can be
triggered from similar redox and hormone imbalances. This is in agreement with the idea exposed by Mittler and associates (2011). In their study, the authors suggest that ROS-mediated signaling possesses a specific signature (probably a set of combinatorial factors acting in parallel) that provides the specificity to trigger different responses.

In conclusion, our findings indicate that a constitutive expression of TMV CP and MP triggers specific stress and defense-like phenotypes and modulates defense responses in a complex manner. It is evident that the mechanisms underlying disease susceptibility and tolerance or resistance depend on a complex regulatory network, and viruses are able to disrupt these fine tunings. Therefore, in this study, we dissected the effect of TMV CP and MP proteins alone or in combination without considering the timing of expression involved in an infection. MP seems to act as a defense elicitor, altering ROS and SA levels, and it is the viral component responsible for enhancing susceptibility against hemibiotrophic and necrotrophic pathogens. CP seems to play a negative effect on host defense by reducing the expression level of defense transcripts like PR-1 and RDR-1. These individual effects reflect the ability of a virus to control its own virulence, given that extremely high virulence is not advantageous, because it could deplete host resources excessively fast. The more efficient viruses are those which achieved balanced behavior and can coexist with hosts without altering their physiology and development.

MATERIALS AND METHODS

RNA extraction and microarray analysis.

To detect differential gene expression between hybrid MP×CPPT42W- and mp×cpPT42W-silenced tobacco transgenic plants, a microarray analysis was performed using a Tomato 10K Affymetrix GeneChip. Leaf tissues from 6-week-old tobacco plants were collected and pools of three plants were homogenized with a mortar and pestle in liquid nitrogen. Next, total RNA was extracted with an RNeasy plant mini kit (Qiagen, Valencia, CA, U.S.A.). RNA quality was assessed with a BioAnalyzer 2100 (Agilent Technologies), and RNA concentration was determined with a spectrophotometer (Nanodrop ND-1000; NanoDrop Technologies, Wilmington, DE, U.S.A.). Each microarray experiment was repeated three times with independent biological replicates (pools of three plants each) following Affymetrix’s instructions manual. Raw hybridization data were preprocessed with robust multichip average (Irizarry et al. 2003). The statistical analysis of preprocessed matrix data was carried out in two phases. First, an analysis of variance (ANOVA) model was applied to keep sequences showing differences between treatments with \( P < 0.10 \). In this way, the original matrix of 10,209 genes was reduced to 1,667. Then, a selection based on \( P \) values with a cutoff of 0.05 to control FDR yielded 28 genes (Table 1). The statistical analysis was performed using InfoStat Statistical software (InfoStat Group, 2004; National University of Córdoba, Argentina). Raw data of the experiment were deposited at Gene Expression Omnibus (GSE37905).

ROS accumulation and cell death detection in tobacco leaves.

For DAB histochemical staining, leaves from control and transgenic tobacco plants were collected and were placed in 1 liter of DAB-HCl, pH 3.8, at a rate of 1 mg of leaf material per milliliter of DAB-HCl, as previously described (Thordal-Christensen et al. 1997) (Sigma-Aldrich, Munich, Germany). For NBT staining, leaves were submerged in an NBT 0.1% solution in 50 mM potassium phosphate buffer, pH 7.8, as previously described by Wohlgenuth and associates (2002). Solutions were infiltrated into leaf tissues by 2-min vacuum shocks in a vacuum chamber; then, the infiltrated leaf tissues were incubated overnight (for DAB staining) or for 2 h for NBT staining. Leaves were cleared in boiling ethanol (96%, vol/vol) to remove chlorophyll and they were examined under a stereo light microscope. \( \text{H}_2\text{O}_2 \) was visible as a brown precipitate in the tissue and superoxide anion was detected as a blue formazan precipitate. Subsequently, for trypan blue staining, plant leaf tissues were boiled for 10 min in a solution composed of 10 ml of 90% (vol/vol) lactic acid, 10 ml of 87% (vol/vol) glycerol, 10 ml of trypan blue solution, and 10 g of phenol. Leaves were finally cleared with chloral hydrate (2.5 mg/ml\(^{-1}\)), conserved in ethanol 96% (vol/vol) and examined under stereo light microscope.

Real-time qPCR.

Total RNA was isolated from 50 to 75 mg of frozen tobacco leave tissues using Trizol Reagent (Invitrogen), quantified (NanoDrop Technologies), and treated with DNase I (Invitrogen, Carlsbad, CA, U.S.A.). First-strand cDNA was synthesized using MMLV (Invitrogen) and random primers according to the manufacturer’s instructions (Invitrogen). The oligonucleotide primer sets used for real-time qPCR analysis were designed using Primer Express 2.0 software (Applied Biosystems, Foster City, CA, U.S.A.). Experiments were carried out using four or five biological replicates in an Applied Biosystems 7500 equipment (the experimental conditions used following MIQE requirements are provided in Supplementary Table S1). \( N. tabacum \) elongation factor-1\( \alpha \) (EF1\( \alpha \) SGN-U446573) was used as an internal control after the evaluation of stability of four reference candidate genes (Supplementary Fig. S3). All primer lists are set in Supplementary Table S2. RT-qPCR data analysis and primer efficiencies were obtained using LinRegPCR software (Remakers et al. 2003). A reference gene was used to standardize the expression of a given target gene; then, a ratio between treatments was calculated using the algorithm developed by Pfaffl and associates (2002). Relative expression ratios and statistical analysis were performed using fgStatistics software interface (J. A. Di Rienzo, personal communications). The cut-off for statistically significant differences was set as *, **, and *** indicate \( P \) value < 0.05, 0.01, and 0.005, respectively.

ASC/DHA measurement.

The method used to determine the amount of total (ASC) and oxidized (DHA) ascorbic acid uses high-performance liquid chromatography (HPLC) with amperometric electrochemical detection and it has been essentially described in a previous work (Diliberto et al. 1983). We made the following minor modifications to this protocol. Detection of ascorbic acid was achieved by injecting samples onto a silica-based, reversed-phase C\( \_18 \) column (particle size 5 pm, 150 x 4.6 mm, HPLC-5s, Bio-Sil; Bio-Rad, Munich). The mobile phase consisted of a K\( \text{H}_2\text{PO}_4 \) buffer (100 mM) at \( \text{pH} \) 3.0 (with phosphoric acid) and was delivered isocratically at a flow rate of 0.5 ml/min. Ascorbic acid resulted in a peak at 3.5 min. Total ASC (reduced plus oxidized) concentration was determined after reduction with dithiothreitol (1 mM). The amount of oxidized DHA was then estimated as the difference in peak area between unreduced and reduced samples.

SA and JA determination.

Extraction and purification procedure. SA and JA were extracted from lyophilized leave tissues according to the protocol of Durbanshi and associates (2005), with some modifications. For JA quantification, 100 ng of JA (2 H\(_6\))JA and 12-oxophytodienoic acid (2H\(_5\)) OPDA were used as internal stand-
ardrs. For SA quantification, the internal standard used was 100 ng of SA-(2H6).

Chromatography and mass spectrometry procedures (liquid chromatography electrospray with negative ionization tandem mass spectrometry). HPLC was performed using a Waters (Milford, MA, U.S.A.) Alliance 2690 system. Aliquots (20 μl) were injected on a Nucleosil ODS reversed-phase column C18 (100 by 2.1 mm, 3 μm). Phytohormones were eluted using an initial gradient of 40% methanol and 60% water/ethanol and, 25 min later, with a gradient of 80% methanol and 20% water:ethanol. Identification and quantification were achieved introducing effluents with an electrospray with negative ionization tandem mass spectrometry into a triple-quadrupole mass spectrometer (Quattro Ultima; Micromass, Manchester, U.K.) from coupled HPLC. For identification, we used mass spectrometry operated with multiple reaction monitoring mode and MassLynx v.4.1 software (Micromass) and, for quantification, we used QuanLynx v.4.1 software (Micromass).

TMV infection procedure.

For inoculation, a single leaf of each plant was dusted with Carborundum and 20 μl of semipurified TMV virus diluted in 20 mM NaHPO4 (pH 7) was added; then, the surface of the Carborundum and 20 TMV infection procedure.

Chromatography and mass spectrometry procedures (liquid chromatography electrospray with negative ionization tandem mass spectrometry). HPLC was performed using a Waters (Milford, MA, U.S.A.) Alliance 2690 system. Aliquots (20 μl) were injected on a Nucleosil ODS reversed-phase column C18 (100 by 2.1 mm, 3 μm). Phytohormones were eluted using an initial gradient of 40% methanol and 60% water/ethanol and, 25 min later, with a gradient of 80% methanol and 20% water:ethanol. Identification and quantification were achieved introducing effluents with an electrospray with negative ionization tandem mass spectrometry into a triple-quadrupole mass spectrometer (Quattro Ultima; Micromass, Manchester, U.K.) from coupled HPLC. For identification, we used mass spectrometry operated with multiple reaction monitoring mode and MassLynx v.4.1 software (Micromass) and, for quantification, we used QuanLynx v.4.1 software (Micromass).

TMV RNA in vitro transcription and tobacco inoculation.

For TMV RNA infections, cloned DNAs from TMV infectious viral RNAs were linearized with KpnI and in vitro transcribed using a MEGAscript T7 in vitro transcription kit (Ambion, Austin, TX, U.S.A.). m7G (5′)ppp(5′)G Cap Analog (Ambion) was added to the reaction to synthesize 5′ capped RNA molecules. Transcripts were rubberized in one single leaf of 6-week-old tobacco plants as described above. Eight plants from each phenotype (CP T42W, MP, MP×CP T42W, mp×cp T42W*, and SX controls) were used in this experiment and samples (two leaf discs from each plant) were collected at the indicated times. ELISA experiments were performed as described previously (Bazzini et al. 2006).

P. syringae inoculation procedure.

Bacterial strains and infection. P. syringae pv. tabaci strains (a gift from M. Alvarez) were cultured overnight in King’s B medium containing antibiotic selection media rifampicin at 25 mg/liter and tetracycline at 2 mg/liter at 28°C with 200 rpm shaking; subsequently, they were centrifuged. Bacterial pellets were resuspended in 10 mM MgCl2 and adjusted to 10⁶ CFU/ml. Fully expanded leaves of 6-week-old tobacco plants were inoculated with bacteria or 10 mM MgCl2 (mock-inoculated plants) using a needless hypodermic syringe. The experiment was repeated twice with similar results. Bacterial counts. At 48 h postinfection, leaf discs from infiltrated zones were collected in a 10 mM MgCl2 solution. To release bacteria, tissue was homogenized by grinding with a sterile pestle. Protein content was quantified with a Quick Start Bradford protein assay kit (Bio-Rad) and homogenized samples were adjusted to equivalent protein concentration. Then, 20-μl dilutions (concentration adjusted for 20 to 400 colonies/plate) were plated in King’s B selective medium and incubated overnight at 28°C. The bacterial plate count = number of colonies/volume of plates × dilution factor. Statistical analyses were performed with Kruskal-Wallis and Dunns post test and differences were calculated between transgenic plant lines and SX controls.

S. sclerotiorum inoculation procedure.

A wild-type isolate of S. sclerotiorum (a gift from A. Escande and D. Barreto) was grown on potato dextrose agar plates as previously described by Godoy and associates (1990). After 4 days, 5-mm mycelia agar plugs were excised and inverted on the upper surface of 10 tobacco leaves per line. Infected leaves were excised, kept in petri plates containing a wet filter paper to ensure high humidity, and incubated at 26°C under a 24-h light photoperiod. Disease symptoms were observed every 24 h over a period of 6 days. The experiment was repeated twice and data of the representative experiment are shown. Statistical analysis were performed with one-way ANOVA and Dunnett post test by comparing leaf necrotic areas between transgenic plant lines and controls SX.

GMP1 VIGS procedure.

GMP1 cloning procedures and vector construction. To amplify a GMP1 cDNA fragment from N. tabacum, total RNA was isolated and subjected to RT-PCR using forward (5′-GT ACTCGGCTGAGGCCATTG-3′) and reverse (5′-CCACGAGTAAATTCCAGCG-3′) primers. The resulting 506-bp products were cloned into pCR8/GW/TOPO vector (Invitrogen) and verified by sequencing. GMP1 506-bp fragments weresubcloned into a pTRV2-GW (Liu et al. 2002) vector by performing an LR recombination reaction. pTRV2-gmp1 plasmids were electroporated into Agrobacterium sp. strain GV3101 and cells were grown on Luria-Bertani (LB) medium with gentamycin at 50 mg/liter and kanamycin at 50 mg/liter. Agroinfiltration and plant treatments. Agrobacterium GV3101 cultures containing the pTRV1, pTRV2-gfp (control), and pTRV2-gmp1 cassettes were grown overnight at 28°C in LB selective medium. Cells were harvested by centrifugation and resuspended in infiltration media (10 mM MgCl2, 10 mM morpholine ethane sulfonic acid, and 200 mM acetosyringone), adjusted to an optical density at 600 nm of 1, and kept at room temperature for 3 to 4 h. Equivalent aliquots of GV3101-pTRV1 and pTRV2 were mixed immediately before inoculation. N. benthamiana plants from four to five leaves were agroinfiltrated as previously described (Liu et al. 2002). Sample leaves were collected 20 days after infiltration, then immediately frozen in liquid nitrogen and stored at −80°C until GMP1 silencing analysis by RT-qPCR. Primer sequences used were forward (5′-TTCCTAAAATTGCCAGCG-3′) and reverse (5′-CGCAAATCCTG GTCCAATCAA-3′) primers. In all, 15 plants from each genotype (NbTRV-gfp and NbTRV-gmp1) were used at the VIGS experiments.

Statistical analysis.

Statistical analysis were performed with one-way ANOVA test with Dunnett post-test GraphPad Prism 5 or Kruskal-Wallis using InfoStat software (InfoStat version 2008; Grupo InfoStat. FCA, Universidad Nacional de Córdoba, Argentina). The significance level for all post tests was α = 0.05.

ACKNOWLEDGMENTS

We thank J. Di Rienzo for assistance with microarray and RT-qPCR analysis, M. Alvarez from UNC (Córdoba, Argentina), who kindly pro-
vided *Pseudomonas syringae pv. tabaci* isolates, A. Escande and D. Barreto from Instituto Nacional de Tecnología Agropecuaria (INTA, Buenos Aires, Argentina) for the generous gift of *Sclerotinia sclerotiorum* isolates; V. Mongelli, A. Bazzini, and F. Carrari for helpful discussions; and M. del Vas and J. Sabio y García for assistance with English-language editing. This research was supported by PICT 2005 number 32598 from Agencia Nacional de Promoción Científica y Tecnológica (ANPCyT) and by project PE 243532 of the (INTA). G. Conti and M. C. Rodríguez hold a fellowship from CONICET and C. A. Manacorda a fellowship from INTA.

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