MicroRNA396-Targeted SHORT VEGETATIVE PHASE Is Required to Repress Flowering and Is Related to the Development of Abnormal Flower Symptoms by the Phyllody Symptoms1 Effector¹

Chiao-Yin Yang², Yu-Hsin Huang², Chan-Pin Lin², Yen-Yu Lin, Hao-Chun Hsu, Chun-Neng Wang, Li-Yu Daisy Liu, Bing-Nan Shen, and Shih-Shun Lin*

Institute of Biotechnology (C.-Y.Y., Y.-H.H., C.-P.L., Y.-Y.L., B.-N.S., S.-S.L.), Genome and Systems Biology Degree Program (S.-S.L.), Department of Plant Pathology and Microbiology (C.-Y.Y., Y.-H.H., C.-P.L., Y.-Y.L.), Institute of Ecology and Evolutionary Biology, Department of Life Science (H.-C.H., C.-N.W.), and Department of Agronomy (L.-Y.D.L.), National Taiwan University, Taipei 106, Taiwan; and Agricultural Biotechnology Research Center, Academia Sinica, Taipei 115, Taiwan (S.-S.L.)

ORCID IDs: 0000-0002-9951-687X (C.-P.L.); 0000-0002-7295-5004 (S.-S.L.).

Leafy flowers are the major symptoms of peanut witches' broom (PnWB) phytoplasma infection in Catharanthus roseus. The orthologs of the phyllody symptoms1 (PHYL1) effector of PnWB from other species of phytoplasma can trigger the proteasomal degradation of several MADS box transcription factors, resulting in leafy flower formation. In contrast, the flowering negative regulator gene SHORT VEGETATIVE PHÂSE (SVP) was up-regulated in PnWB-infected C. roseus plants, but most microRNA (miRNA) genes had repressed expression. Coincidentally, transgenic Arabidopsis (Arabidopsis thaliana) plants expressing the PHYL1 gene of PnWB (PHYL1 plants), which show leafy flower phenotypes, up-regulate SVP of Arabidopsis (AtSVP) but repress a putative regulatory miRNA of AtSVP, miR396. However, the mechanism by which PHYL1 regulates AtSVP and miR396 is unknown, and the evidence of miR396-mediated AtSVP degradation is lacking. Here, we show that miR396 triggers AtSVP messenger RNA (mRNA) decay using genetic approaches, a reporter assay, and high-throughput degradome profiles. Genetic evidence indicates that PHYL1 plants and atmir396a-1 mutants have higher AtSVP accumulation, whereas the transgenic plants overexpressing MIR396 display lower AtSVP expression. The reporter assay indicated that target-site mutation results in decreasing the miR396-mediated repression efficiency. Moreover, degradome profiles revealed that miR396 triggers AtSVP mRNA decay rather than miRNA-mediated cleavage, implying that AtSVP caused miR396-mediated translation inhibition. We hypothesize that PHYL1 directly or indirectly interferes with miR396-mediated AtSVP mRNA decay and synergizes with other effects (e.g. MADS box transcription factor degradation), resulting in abnormal flower formation. We anticipate our findings to be a starting point for studying the posttranscriptional regulation of PHYL1 effectors in symptom development.

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Phytoplasmas are obligate bacteria without cell walls that infect numerous plants, causing severe agricultural losses (Lee et al., 2000). The typical symptoms on phytoplasma-infected plants include witches' broom, proliferative branches, leaf yellowing, and small leaves (Lee et al., 2000). The flowers of infected plants exhibit sterility, abnormal morphology, virescence, and phyllody (herein referred to as leafy flower; Himeno et al., 2011). Peanut witches' broom (PnWB) phytoplasma-infected *Catharanthus roseus* plants exhibit leafy flower symptoms, which were classified into five stages depending on their severity as described previously (Yang, 1985; Liu et al., 2014). Stage 4 (S4) leafy flowers exhibited leaf vein patterning with phyllody petals and intumescent ovaries (Fig. 1A; Liu et al., 2014).

The draft genome sequence of the PnWB phytoplasma contains 13 chromosomal contigs with 562,473 bp and 421 protein-coding genes (Chung et al., 2013). Chung et al. (2013) suggested that the 48 putative PnWB effectors are similar to known pathogenicity effectors of phytoplasmas. In aster yellows witches' broom (AYWB)

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² These authors contributed equally to the article.

^{*} Address correspondence to linss01@ntu.edu.tw.

The author responsible for distribution of materials integral to the findings presented in this article in accordance with the policy described in the Instructions for Authors (www.plantphysiol.org) is: Shih-Shun Lin (linss01@ntu.edu.tw).

S.-S.L. and C.-P.L. conceived the original screening and research plans; S.-S.L. and C.-P.L. supervised the experiments; C.-Y.Y. and Y.-H.H. performed most of the experiments; Y.-Y.L., H.-C.H., C.-N.W., and B.-N.S. provided technical assistance; L.-Y.D.L. performed the statistical assays; S.-S.L. designed the experiments and analyzed the data; S.-S.L. conceived the project and wrote the article with contributions of all authors.

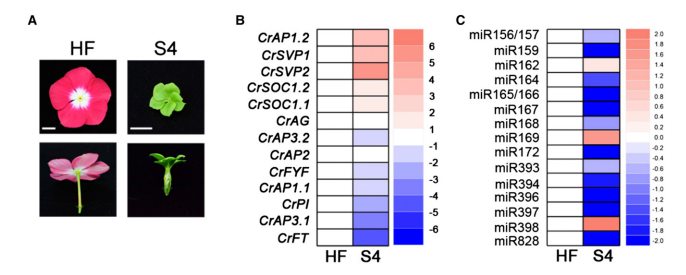


Figure 1. The leafy flower phenotypes of PnWB phytoplasma-infected *C. roseus* plants. A, The healthy flower (HF) and S4 PnWB-infected leafy flower symptoms are presented in the left and right columns, respectively. Bars = 1 cm. B and C, The mRNA (B) and microRNA (miRNA; C) expression of the HF and S4 in *C. roseus* flowers are represented by heat maps. Red indicates up-regulated expression, and blue indicates down-regulated expression. The values at right indicate the results of the log₂ (S4/HF) formula. *CrAG, AGAMOUS* of *C. roseus*; *CrFYF, FOREVER YOUNG FLOWER* of *C. roseus*; *CrPI, PISTILLATA* of *C. roseus*.

phytoplasma, the secreted AYWB PHYTOPLASMA PROTEIN54 effector (SAP54) causes a leafy flower phenotype in transgenic Arabidopsis (*Arabidopsis thaliana*) plants (*SAP54* plants; MacLean et al., 2011). Moreover, the phyllody symptoms1 (PHYL1) effector of onion yellows phytoplasma wild-type line, a SAP54 ortholog, also manifested an identical leafy flower phenotype in *PHYL1*-overexpressing plants (Maejima et al., 2014).

PHYL1 and SAP54 interact with MADS domain transcription factors (MTFs) in Arabidopsis, such as SEPALLATA3 (AtSEP3), APETALA1 (AtAP1), and CAULIFLOWER (AtCAL; MacLean et al., 2014; Maejima et al., 2014). These MTFs are key regulators in floral development and were triggered by SAP54/ PHYL1-mediated ubiquitin/26S proteasomal degradation by interacting with the RADIATION SENSITIVE23 (RAD23) family, resulting in interference in floral homeotic gene expression levels (MacLean et al., 2014; Maejima et al., 2014). Consistently, the ap1 mutant, ap1/cal double mutant, and sep1/sep2/sep3/sep4 quadruple mutant manifest various types of leafy flower (e.g. leaf-like sepals, secondary flowers, cauliflower-like inflorescences, or loss of floral meristem determinacy; Bowman et al., 1993; Ditta et al., 2004; Gregis et al., 2006). These results suggest that phytoplasma SAP54/PHYL1 is a key effector for triggering positive regulators of MTF degradation to switch the phase transition of plants from the reproductive stage to the vegetative stage.

In PnWB-infected *C. roseus* plants, the expression levels of *SHORT VEGETATIVE PHASE1* (*CrSVP1*) and *CrSVP2* were up-regulated in S4 leafy flowers (Liu et al., 2014). The SVP of Arabidopsis (AtSVP) is an MTF and functions as a negative regulator of floral development as part of a complex with FLOWERING LOCUS C (AtFLC)

to control flowering time by negatively regulating FLOWERING LOCUS T (AtFT) and SUPPRESSOR OF OVEREXPRESSION OF CONSTANS1 (AtSOC1) by directly binding their promoter sequences (Lee et al., 2007b; Li et al., 2008; Jang et al., 2009; Irish, 2010). Moreover, AtSVP also interacts with other components, such as AP1 and AGAMOUS-LIKE24 (AGL24), during stages 1 and 2 of flower development to repress floral homeotic genes that control petal, stamen, and carpel identity (Gregis et al., 2009). The svp mutant (svp32) of Arabidopsis exhibits an early-flowering phenotype, whereas constitutive expression of AtSVP in Arabidopsis delays flowering (Hartmann et al., 2000; Gregis et al., 2006; Lee et al., 2007a). Moreover, transgenic Arabidopsis expressing AtSVP orthologs from Actinidia spp. or rice (Oryza sativa) showed leaf-like sepals and secondary flower phenotypes (Trevaskis et al., 2007; Wu et al., 2012). In addition to MTF down-regulation, phytoplasma might induce leafy flower symptom development via upregulated SVP expression.

MTF plays a role in the regulation of gene expression, whereas miRNA plays a role in posttranscriptional regulation. Whether phytoplasma effectors alter the miRNA expression levels for symptom development is an interesting question. MiRNAs regulate numerous transcription factors involved in plant development through RNA cleavage or translation inhibition (Kidner and Martienssen, 2005; Voinnet, 2009). In contrast to animal miRNAs, plant miRNAs display a high degree of sequence complementarity to their target mRNAs and guide cleavage of the target RNA via ARGONATUTE1 (AGO1; Kidner and Martienssen, 2005; Voinnet, 2009). In animals, the imperfect miRNA/target pairwise region results in translation inhibition and, consequently, decay

of the target mRNA (Bazzini et al., 2012; Djuranovic et al., 2012). However, the mechanism of translation inhibition in plants is not well known.

Chorostecki et al. (2012) proposed a predicted miR396 target site on the MADS box region of the AtSVP mRNA. Nevertheless, the lack of direct evidence from 5' RACE data left them unable to demonstrate miR396-mediated AtSVP cleavage (Chorostecki et al., 2012). In contrast, the miR396-mediated GROWTH-REGULATING FACTOR (AtGRF) family of Arabidopsis features a bulge between the seventh and eighth positions of miR396 that is highly conserved among different plant species. The 5' RACE data indicated that the bulge structure did not affect miR396-mediated AtGRF cleavage. However, interaction along the 5' portion of the miRNA is the feature most relevant to its activity and cleavage efficiency (Mallory et al., 2004; Schwab et al., 2005; Lin et al., 2009), and modifying the pairing status of the 5' portion of miR396 through mutagenesis alters the efficiency of AtGRF2 cleavage (Debernardi et al., 2012). Moreover, Li et al. (2013) demonstrated that a 5' miRNA mismatch triggers miRNA-mediated target RNA translation inhibition in Arabidopsis. These results suggest that mispairing of the 5' portion of the miRNA plays a role in modulating cleavage efficiency and translation inhibition during target gene regulation.

We previously identified miR396 of *C. roseus* via whole-transcriptome analysis and small RNA profiling by next-generation sequencing (Chang, 2012; Liu et al., 2014). There are two mismatches in the 5' portion of the miR396 (fourth and fifth positions) on the pairwise region of miR396/*CrSVPs* (Chang, 2012). Therefore, this study seeks to answer several questions: (1) whether *CrSVP1*/2 are targets of miR396; (2) whether the regulation approaches differ between miR396/*CrSVPs* and miR396/*CrGRFs* due to the two mismatches in the 5' portion of the miRNA; (3) whether the PHYL1 effector of PnWB also determines leafy flower phenotype in Arabidopsis; and (4) whether PHYL1 alters MTF and miRNA expression levels.

In this study, transgenic Arabidopsis expressing the *PHYL1* gene of PnWB (*PHYL1* plant) was generated and showed an identical leafy flower phenotype. The leafy flowers of *PHYL1* plants exhibited a lower miR396 expression level but higher *AtSVP* and *AtGRF1* expression levels compared with normal flowers of Columbia-0 (Col-0) plants. Moreover, we demonstrated that miR396 triggers *SVP* down-regulation via mRNA decay rather than cleavage based on degradome analysis. The mechanisms of PHYL1-altered miR396/*SVP* expression levels and miR396-mediated *SVP* down-regulation were investigated.

RESULTS

Abnormal Gene and miRNA Expression in PnWB-Infected C. roseus

Deep sequencing was used to compare the whole transcriptome with the small RNA profiles of *C. roseus*

HF and S4 PnWB-infected leafy flowers (Fig. 1A). The floral homeotic gene expression levels were compared between HF and S4 flowers. The expression levels of CrFT, PISTILLATA of C. roseus, FOREVER YOUNG FLOWER of C. roseus, CrAP1.1, CrAP3.1, and CrAP3.2, but not CrAP1.2, CrSOC1.1, and CrSOC1.2, which enhance flowering, were repressed in the S4 flowers (Fig. 1B). In contrast, the expression levels of CrSVP1 and CrSVP2 were increased in S4 flowers (Fig. 1B). AGAMOUS of C. roseus and CrAP2 were not differentially expressed between HF and S4 flowers (Fig. 1B). In addition, we observed that most *C. roseus* miRNAs were significantly down-regulated in the S4 flowers, with only the upregulation of miR162, miR169, and miR398 observed (Fig. 1C). These results suggest that PnWB effectors or other bacterial proteins might directly or indirectly alter either the miRNA gene expression profile of C. roseus plants (CrMIR genes) or the floral homeotic gene expression levels in PnWB-infected plants.

The PHYL1 Effector of PnWB Causes Leafy and Abnormal Flower Phenotypes and Represses miR396 Expression in Arabidopsis

The SAP54 effector of AYWB causes leafy flower in transgenic Arabidopsis (MacLean et al., 2011). We identified a PHYL1 effector gene in the PnWB genome sequence (Chung et al., 2013) that shares 61.3% amino acid sequence similarity with AYWB and 56% similarity with PHYL1 of the onion yellows phytoplasma wildtype line (Fig. 2A) and determined whether PHYL1 of PnWB can cause leafy flower and alter miRNA expression. PHYL1 plants, which overexpress the PHYL1 of PnWB, exhibit two types of leafy flowers (Fig. 2B) and three types of abnormal flowers (Fig. 2C), depending on the severity of the abnormal phenotype in the flowers. The type I leafy flower phenotype exhibits sepal enlargement, petal shrinking, and stigma elongation (Fig. 2B). The type II leafy flower phenotype features a secondary flower growing out from the primary flower (Fig. 2B). In some cases, a cauliflower-like inflorescence with narrow interspaces and short pedicles results (Fig. 2B). Three types of abnormal flowers include asymmetric petals, abnormal petal, and stamen numbers (five to six petals and seven to eight stamens; AF-I), swollen stigma, and secondary flower growth (Fig. 2C). We assume that various phenotypes occurred on PHYL1 plants because of the various expression levels of PHYL1.

Interestingly, in *PHYL1* and *SAP54* plants, most miRNAs were up-regulated compared with Col-0 plants, whereas miR156h, miR159, miR390, and miR396 were repressed, indicating that both effectors have similar functions in *CrMIR* gene regulation (Fig. 3A). However, PnWB infection inhibits most *CrMIR* genes (Fig. 1C). We hypothesize that the other effectors of PnWB might be involved in the repression of *CrMIR* genes. Furthermore, PHYL1 and SAP54 have specificity in the repression of particular miRNAs. Indeed, the northern-blot data also

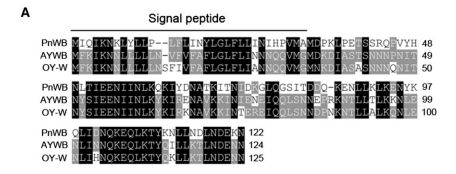
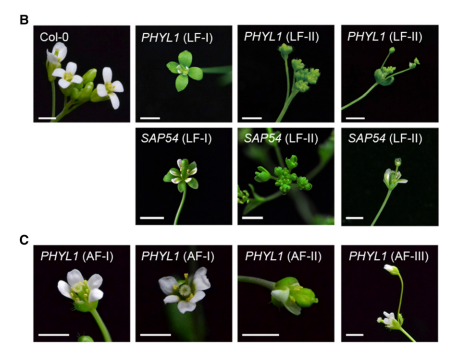


Figure 2. The phytoplasma effectors trigger the leafy flower phenotype. A, Alignment of orthologous amino acids of PHYL1. OY-W, Onion yellows phytoplasma wild-type line. B, Leafy flower phenotypes of *PHYL1* and *SAP54* plants. The type I leafy flower is indicated as LF-I, and the type II leafy flower is indicated as LF-II. Bars = 0.5 cm. C, The abnormal flower phenotype of *PHYL1* plants. The type I to III phenotypes are indicated as AF-I, AF-II, and AF-III, respectively. Bars = 0.5 cm.



indicated that the leafy flowers of *PHYL1* plants exhibited lower miR396 accumulation (0.57-fold) compared with Col-0 plants (Fig. 3B).

Identification of the MIR396 Gene of C. roseus

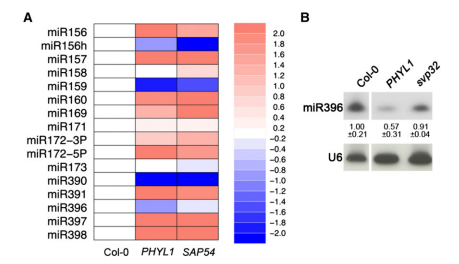
The hairpin structures of the miR396 precursor of C. roseus (DDS44007) and its miRNA/miRNA* pairing positions are shown in Figure 4A. The miR396/ miR396* duplex exhibits a two-nucleotide overhang at the 3' end (Fig. 4A). Remarkably, 14,256 reads of the mature form of miR396 were obtained for the HF flowers, whereas only 1,482 reads of miR396* were obtained (Fig. 4A). These results satisfy the general rule that less miRNA* than the mature form of the miRNA is required in vivo because the miRNA* exhibits a rapid turnover rate due to cleavage by AGO1 (Guo and Lu, 2010). Importantly, the amount of miR396 was reduced (2,587 reds) in S4 flowers, which is consistent with previous observations (Fig. 4A). In addition, the complementary DNA (cDNA) of the miR396 precursor was cloned from the C. roseus flower by reverse transcription (RT)-PCR for further investigation (data not shown), suggesting that the miR396 precursor exists in *C. roseus* plants.

The MADS Box Domain of the SVP Gene Contains a Potential miR396 Target Site

Based on sequence pairing, a putative miR396 target site was identified on the MADS box domain of the *SVPs* (Fig. 4B); however, the miR396-mediated cleavage of *SVP* was not confirmed by 5' RACE (Chorostecki et al., 2012). The target sequences in *CrSVP1* and *CrSVP2* were highly conserved with that of *AtSVP* (AT2G22540) of Arabidopsis, and four mismatches (the fourth, fifth, 19th, and 20th positions of miR396) were identified in the pairwise region of miR396/*CrSVP1* or miR396/*AtSVP* (Fig. 4B). However, the pairwise region of miR396/*CrSVP2* had an extra G-to-A mismatch at the seventh position of miR396 (Fig. 4B).

MiR396 was down-regulated 0.06-fold in the S4 leafy flower compared with the HF (Fig. 4C). Moreover, semi-RT-PCR results indicated that the levels of

Figure 3. Phytoplasma effectors interfere with miRNA expression. A, MiRNA expression in flowers of Col-0, *PHYL1*, and *SAP54* plants as represented by a heat map. Red indicates upregulated expression, and blue indicates downregulated expression. The values at right indicate the results of the log₂ (Col-0/*PHYL1* or Col-0/*SAP54*) formula. B, MiR396 detection in Col-0, *PHYL1*, and *svp32* plants by northern blot. U6 was used as a loading control. The numbers indicate the relative expression of miR396 compared with Col-0 plants.



CrSVP1 were up-regulated 3.2- and 3.5-fold in the S4 leafy flower and healthy leaf samples, respectively, compared with the HF (Fig. 4D). In addition, *CrSVP2* was up-regulated 3.4- and 4.6-fold in S4 and healthy leaf, respectively (Fig. 4D). These data suggest that PnWB infection alters the expression of *CrSVP1/2* and miR396.

SVPs Contain Mismatch Pairings at the Target Site in the 5' Portion of miR396

Alignment of the *SVP* genes revealed highly conserved miR396 target sites in various species. Fewer than five mismatches were observed on these *SVP*-miR396 pairwise regions, whereas 94% of MADS box genes had more than six mismatches spread throughout parallel regions of *AtSVP*, such as *AtAGL24* and *AtFLC* (Fig. 5A; Supplemental Fig. S1). The *AtGRF* gene family is also the target of miR396 (Jones-Rhoades and Bartel, 2004; Wang et al., 2004; Debernardi et al., 2012). However, the *AtGRF*-miR396 pairwise region exhibits mostly complementary pairing that differs from the *SVP*-miR396 pairing (Fig. 5B). Thus, we assume that the miR396-mediated degradation efficiency or approaches between miR396/*SVPs* and miR396/*AtGRFs* might differ (Fig. 5, A and B).

MiR396-Mediated Down-Regulation of SVPs by Transient Expression

Different concentrations of agrobacteria carrying the MIR396a gene of Arabidopsis (AtMIR396a) were transiently coexpressed with AtSVP in Nicotiana benthamiana plants and were then analyzed via agroinfiltration to evaluate whether miR396 can down-regulate AtSVP. The AtSVP gene was expressed using a Cauliflower mosaic virus 35S promoter to avoid the side effects of endogenous transcription factors that are regulated by miR396. The AtSVP RNA levels were decreased

by increasing miR396 expression levels (Fig. 5C). Similar results were observed for the miR396-mediated down-regulation of *CrSVP1*, *CrSVP2*, and *AtGRF1* (Supplemental Fig. S2). No *CrSVP1* down-regulation occurred when amiRGFP4 was present (an artificial miRNA that targets the *GFP* gene used as a negative control; Supplemental Fig. S3). These data indicate that miR396 directly triggers *SVP* posttranscriptional down-regulation.

To further demonstrate the target gene downregulation by miR396, a reporter assay was used in this study. A DNA fragment (129 nucleotides in length) that contains a target site for AtSVP and AtGRF1 was fused at the 5' end of the YELLOW FLUORESCENT PROTEIN (YFP) gene to generate SVP_{129} -YFP and $GRF1_{129}$ -YFP constructs, respectively (Fig. 6A). Moreover, SVP_{129m} -YFP is a mutant construct that has several mutations at the target site, resulting in eight mismatches in the pairwise region of miR396 (Fig. 6A). The SVP_{129} -YFP and $GRF1_{129}$ -YFP constructs resulted in an approximately 85% reduction of YFP fluorescence when these constructs were coexpressed with miR396, whereas no significant differences were observed regarding YFP fluorescence in the YFP negative control with/without miR396 (Fig. 6, B and C). In contrast, the SVP_{129m}-YFP construct resulted in an approximately 40% reduction of YFP fluorescence when coexpressed with miR396 (Fig. 6, B and C). Notably, the fluorescent spots are present in nuclei (Fig. 6B). The reporter assay demonstrated that miR396 specifically down-regulated AtSVP and AtGRF1.

Flowering Phenotypes of miR396 in Loss and Gain of Function

We used genetic approaches to examine gain of function and loss of function in miR396-mediated *SVP* regulation. A transfer DNA (T-DNA) insertion mutant line, SALK064047, was demonstrated to be an *AtMIR396a* mutant (*atmir396a-1*) in Arabidopsis (Bao

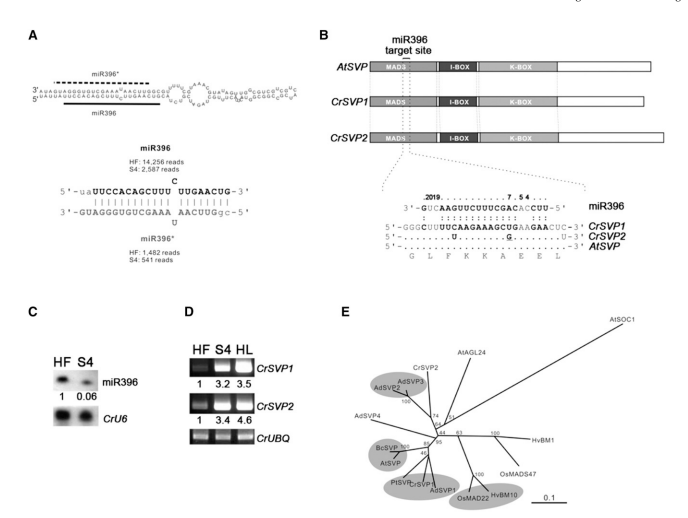


Figure 4. Characterization of MIR396, CrSVP1, and CrSVP2 in C. roseus. A, Illustration of the hairpin structures of the MIR396 precursors of C. roseus and their paired sequences. The mature miR396 (solid line) and miR396* (dashed line) are indicated on the precursor structure (top). The pairing of the miR396 (black) and miR396* (gray) sequences are indicated at bottom. The numbers indicate the deep-sequence reads of miR396 or miR396*. B, Predicted targets of miR396 in the MADS box domain of SVP genes. The nucleotide sequences of the CrSVP1 (DDS_51524), CrSVP2 (DDS_57797), and AtSVP (AT2G22540) targets were aligned, and the paired regions with miR396 are indicated. In the SVP alignment, dots indicate identical nucleotides, whereas boldface letters indicate the nucleotides that are perfectly paired between SVP and miR396. The translated SVP amino acids are presented below the alignment. C, Analysis of miR396 expression between HF and S4 PnWB-infected leafy flower by small RNA northern blot. The small RNA loading quantity was adjusted to 200 ng using the 2100 Bioanalyzer (Agilent). The numbers below indicate the relative fold change in the expression of miR396 regarding the S4 flowers relative to the HFs. CrU6 small nuclear RNA was used as a loading control. D, Semi-RT-PCR analyses of the CrSVP1 and CrSVP2 expression levels in an HF, S4 flower, and healthy leaf (HL). UBIQUITIN (UBQ) of C. roseus was used as an internal control. The numbers indicate the fold change in CrSVP1 and CrSVP2 expression in the S4 flower and HL samples relative to the HF. E, Phylogenetic tree of SVP proteins. The genes sharing greater than 80% amino acid similarity are classified and shaded in gray. AtSOC1 is referred to as an outgroup. The numbers indicate the percentage of 1,000 bootstrap replicates at the appropriate nodes. The scale bar indicates a 0.1 divergence of amino acid substitutions per site.

et al., 2014). T-DNA was inserted at the -450-bp promoter region of *AtMIR396a* (AT2G10606; Fig. 7A). The homozygous *atmir396a-1* plants exhibited only 0.6-fold miR396 expression levels compared with Col-0 plants (Fig. 7B). We assumed that this expression represents a background signal of miR396b in *atmir396a-1* plants (Fig. 7B). The *atmir396a-1* plants exhibit normal flower and flowering time phenotypes, whereas the *svp32* plants display early flowering (Fig. 7, C and D). In addition,

atmir396a-1 plants exhibit 0.5- and 0.3-fold increased AtSVP and AtGRF1 expression, respectively, compared with Col-0 plants (Fig. 7E), indicating that lower miR396 levels induce up-regulation of AtSVP and AtGRF1.

In contrast, transgenic Arabidopsis expressing *CrMIR396* (*CrMIR396* plant) exhibits abnormal flowers with curved sepals and petals and enlarged stigmas, consistent with transgenic Arabidopsis expressing

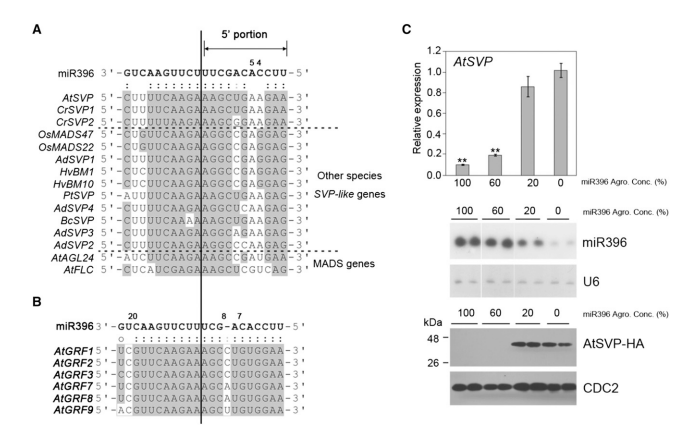


Figure 5. *SVP* contains variation pairing of the 5' portion of miR396 with its target genes. A, Highly conserved miR396 target sites on the *SVP* genes of various species. The alignment of the miR396 target sites in *SVP* genes of various species is shown. The GenBank accession numbers of the *SVP* genes are as follows: *AtSVP* (NM127820), *AtAGL24* (NM118587), *AdSVP1* (JF838212), *AdSVP2* (JF838213), *AdSVP3* (JF838214), *AdSVP4* (JF838215), *HvBM1* (AJ249142), *HvBM10* (EF043040), *OsMADS22* (AB107957), *OsMADS47* (AY345221), *BcSVP* (DQ922945), and *PtSVP* (FJ373210). *AtAGL24* and *AtFLC* (AT5G10140) were chosen as representative *MADS* genes for target-site comparison. The *SVP* nucleotides that pair with the miR396 sequences are indicated in gray. B, The miR396 target-site alignment for *GRFs* (*AtGRF1*, AT2G22840; *AtGRF2*, AT4G37740; *AtGRF3*, AT2G36400; *AtGRF7*, AT5G53660; *AtGRF8*, At4G24150; and *AtGRF9*, AT2G45480). Gray shading indicates the nucleotides of *AtGRFs* paired with the miR396 sequences. C, In vivo analysis of miR396-mediated *AtSVP-HEMAGGLUTININ* (*HA*) efficiency by transient expression. The relative expression levels of *AtSVP* were normalized to that of *NbACTIN* via real-time RT-PCR. Error bars represent set (*n* = 6). The relative expression levels were significantly different from those in Col-0 plants for each RNA sample, based on Student's *t* test: **, *P* < 0.01. The various percentages of miR396 were determined by a small RNA northern blot. U6 small nuclear RNAs were used as loading controls. The various percentages of miR396 are indicated below the real-time RT-PCR and northern-blot data. The AtSVP-HA protein level was detected by western blotting with a 1:10,000 dilution of HA antibody. *CELL DIVISION CONTROL2* (*CDC2*) was used as a protein loading control.

AtMIR396a (AtMIR396a plant; Fig. 7C; Liang et al., 2014). Two individual CrMIR396 plants (lines 8 and 27) exhibited 4.6- and 3.5-fold increased miR396 expression compared with Col-0 plants. Furthermore, AtMIR396a plants exhibited a 1.9-fold increase in miR396 expression (Fig. 7B). CrMIR396 and AtMIR396a plants exhibited late-flowering phenotypes that were correlated with the expression levels of miR396 (Fig. 7D). Moreover, AtSVP and AtGRF1 were down-regulated in CrMIR396 and AtMIR396a plants (Fig. 7E). AtGRF1 was down-regulated by more than 80% by the overexpression of miR396, while AtSVP was downregulated by only 20% to 40% in MIR396 plants (Fig. 7E). These results indicated that CrMIR396 can generate miR396 and has a genetic function similar to that of AtMIR396a in Arabidopsis plants. Moreover,

miR396 regulates *AtGRF1* more efficiently than *AtSVP*.

SVP Orthologs from Different Species Cause Various Flower Phenotypes

Two orthologous *SVP* genes (*CrSVP1* and *CrSVP2*) from *C. roseus* were overexpressed in Arabidopsis (*CrSVP1* and *CrSVP2* plants), and transgene overexpression was confirmed by RT-PCR (Supplemental Fig. S4). However, both plants exhibited normal flower phenotype and flowering time (Fig. 7, C and D). In addition, transgenic Arabidopsis expressing the *SVP2* gene of *Actinidia* spp. (*AdSVP2* plants) exhibited enlarged sepals, small green petals, and a curving pistil

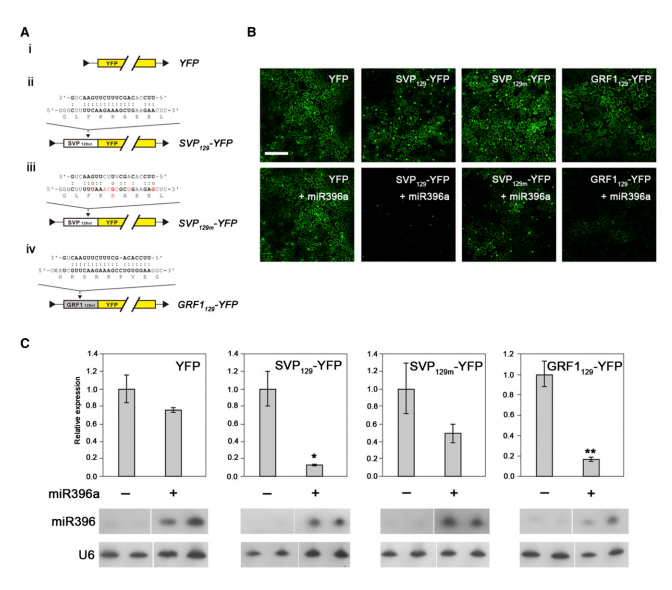


Figure 6. The reporter assay for miR396-mediated targeted down-regulation. A, Schematic of miR396 and its target-site pairing on the *YFP* gene. All of the constructs were constructed on a binary vector under the control of a 35S promoter. A *YFP*-only construct (i) was used as a negative control. The SVP_{129} -YFP construct (ii) represents 129 nucleotides of the 5' end of the SVP gene containing the miR396 target site fused with the YFP gene. The SVP_{129m} -YFP construct (iii) represents 129 nucleotides of the 5' end of the SVP gene containing a mutated miR396 target site fused with the YFP gene. The $GRF1_{129}$ -YFP construct (iv) represents 129 nucleotides of the 5' end of the GRF1 gene containing the miR396 target site fused with the YFP gene. B, The YFP reporter assay was analyzed using confocal microscopy. Bar = 100 μ m. C, Relative average YFP expression levels (log₂) of the reporter assay as determined by confocal microscopy (n = 3). Relative YFP expression levels were significantly different from that of the negative control construct without miR396 present in each sample, based on Student's t test: **, t < 0.01; and *, t < 0.05. MiR396 was detected by northern blot for each sample (two replicates). U6 small nuclear RNA was used as a loading control.

mimicking the leafy flower phenotype of *PHYL1* plants (Fig. 7C); however, the flowering time of the *AdSVP2* plant was similar to that of Col-0 plants (Fig. 7D). In contrast, the *AtSVP* plants showed asymmetric petals of an abnormal flower phenotype identical to *PHYL1* plants (Figs. 2C and 7C). In some cases, the severely abnormal flowers of *AtSVP* plants had five petals with eight stamens (Fig. 7C). These data suggested that the up-regulation of *AtSVP* causes an abnormal flower phenotype. The abnormal flower

phenotype in *AtSVP* plants is consistent with the AF-I of PHYL1 plants, which have five to six asymmetric petals and seven to eight stamens (Figs. 2C and 7C).

Two individual lines of *AtSVP* plants (lines 1 and 2) exhibited late flowering time with 16 to 19 rosette leaves. Indeed, phylogenetic analysis revealed that AtSVP, AdSVP2, CrSVP1, and CrSVP2 are in different groups based on 80% amino acid similarity (Fig. 4E). Moreover, the C-terminal regions of these SVPs differ (Supplemental Fig. S5). These results imply that the

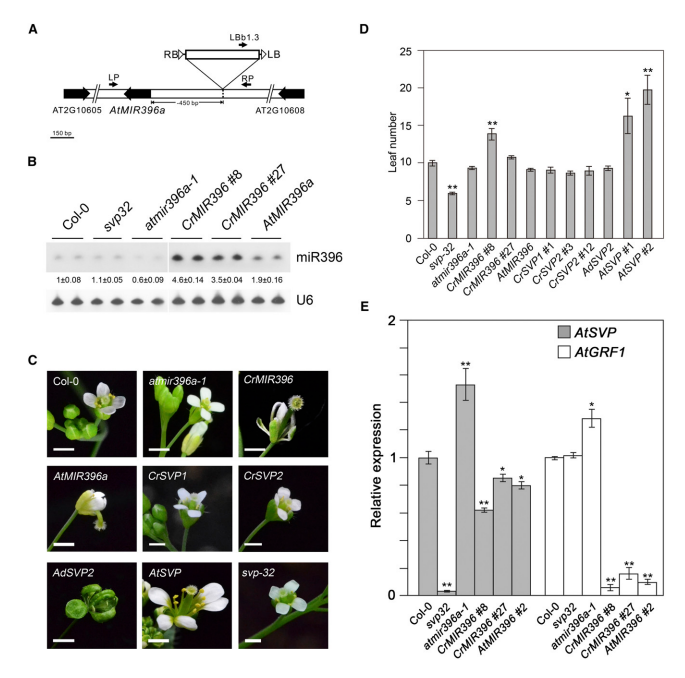


Figure 7. Characterization of miR396 and *SVPs* in Arabidopsis plants. A, Diagram of the *atmir396a-1* mutant (SALK064047). The black arrows indicate the gene regions for AT2G10605, *AtMIR396a* (AT2G10606), and AT2G10608 and their translation directions. The T-DNA insertion site is located at the promoter region (450 bp upstream) of *AtMIR396a*. The left border (LB) and right border (RB) are indicated at the two borders of the T-DNA. The primers LP, RP, and LBb1.3 were used for genotyping via PCR. Bar = 150 bp. B, Analysis of miR396 by small RNA northern blot in Col-0, *svp32* mutant, *atmir396a-1* mutant, two independent *CrMIR396* lines (lines 8 and 27), and *AtMIR396a*-overexpressing plants. U6 small RNA was used as a loading control. The numbers below indicate fold changes in miR396 expression relative to the Col-0 plant. C, *CrSVP1*, *CrSVP2*, *atmir396a-1*, and *svp32* (SALK072930C) mutant plants exhibited normal flower phenotypes. *CrMIR396* and *AtMIR396a* plants exhibited abnormal flower phenotypes. Plants with *AdSVP2* or *AtSVP* overexpression exhibited leafy or abnormal flower phenotypes, respectively. Bars = 0.1 cm. D, Evaluation of the flowering time of various Arabidopsis mutant or transgenic plants. Leaf number represents flowering time. Error bars represent set (n = 20). E, MiR396-mediated *AtSVP* and *AtGRF1* regulation. *AtSVP* and *AtGRF1* were detected by real-time RT-PCR. Relative expression levels were normalized to the level of *AtUBQ*. Error bars represent set (n = 3). Relative expression levels were significantly different from those of the Col-0 plants in each RNA sample, based on Student's *t* test: *, P < 0.05; and **, P < 0.01.

SVP genes in other species have functions regarding flower phenotypes and flowering times that differ from those in Arabidopsis. In addition, the abnormal flower phenotypes in *AdSVP2* and *AtSVP* plants suggest that overexpression of *SVP* may alter flower development. To confirm this hypothesis, *AtSVP* expression was detected in *PHYL1* plants.

SVP Is Up-Regulated in the Sepal and Petals of PHYL1 Plants

Because miR396 expression was reduced in *PHYL1* plants, we evaluated *AtSVP* and *AtGRF1* expression in transgenic plants. Real-time RT-PCR analysis revealed that *AtSVP* and *AtGRF1* are up-regulated in the *PHYL1* leafy flower (Fig. 8A). Moreover, in situ hybridization data indicate strong accumulation of *AtSVP* signals in the leafy flower tissue of *PHYL1* plants, particularly in the basal part of the petals (Fig. 8B, image ii), whereas few *AtSVP* signals were observed in the sepals of Col-0 plants (Fig. 8B, image i). The sense probe for *AtSVP* was used as a negative control for in situ hybridization, and no signal was observed in flower buds of *PHYL1* or Col-0 plants (Fig. 8B, images iii and iv). These results support the hypothesis of miR396-mediated *AtSVP* expression in flower tissue.

MiR396 Triggers SVP Decay

The pairwise fourth and fifth positions are reportedly critical for the efficiency of low miRNA-guided cleavage of the target RNA (Lin et al., 2009). We assumed that the two mismatches at the fourth and fifth positions of the pairwise region will yield differences in cleavage between miR396/AtSVP and miR396/AtGRF1. To test this

hypothesis, a high-throughput degradome approach was used to analyze the mRNA degradation products. Overexpression of miR396 yielded 22,705 degradome reads of degraded 3' AtGRF1 fragments in the CrMIR396 plant, whereas only 800 degradome reads were obtained from Col-0 plants (Fig. 9A). In addition, 3,744 degradome reads of AtGRF1 were obtained from EXORIBONUCLEASE4 mutant (xrn4-5), a 5'-to-3' exoribonuclease mutant, whereas approximately 500 degradome reads were detected in SAP54 plants (Fig. 9A). These data indicate that miR396-mediated AtGRF1 cleavage is dependent on miR396 dosage and that the cleaved 3' AtGRF1 fragment is highly accumulated in xrn4-5 mutants. A 3- to 5-fold increase in AtGRF1 degradome reads was detected in CrMIR396 and xrn4-5 plants compared with Col-0 plants (Fig. 9B). In addition, the other AtGRF targets, Arabidopsis BASIC HELIX-LOOP-HELIX74 (AT1G10120) and AtMMG4.7 (AT5G43060), which are also targets of miR396, also produced significant degradome reads on the corrected target site (Supplemental Fig. S6). These results indicate that miR396 mediated AtGRF1 cleavage at the target site.

While no degradome reads were detected for the target site of *AtSVP* in these plants, approximately 280 various degradome reads were identified at the rear position of the miR396 target site in *CrMIR396* plants, suggesting the presence of miR396-mediated *AtSVP* decay (Fig. 9C, white arrowheads). Total degradome reads of *AtSVP* were increased 5-fold in *CrMIR396* plants compared with Col-0 plants, but no significant reads were detected in either *xrn4-5* or *SAP54* plants (Fig. 9C). Identical results were obtained for the degradome profile of *C. roseus* flowers (Fig. 9E, images i and ii). In total, 52 and 21 degradome reads were detected for the rear positions of the target sites on

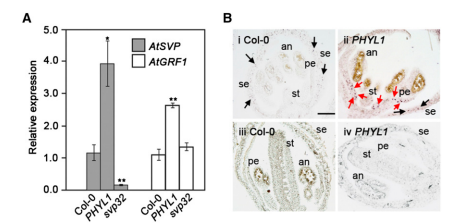
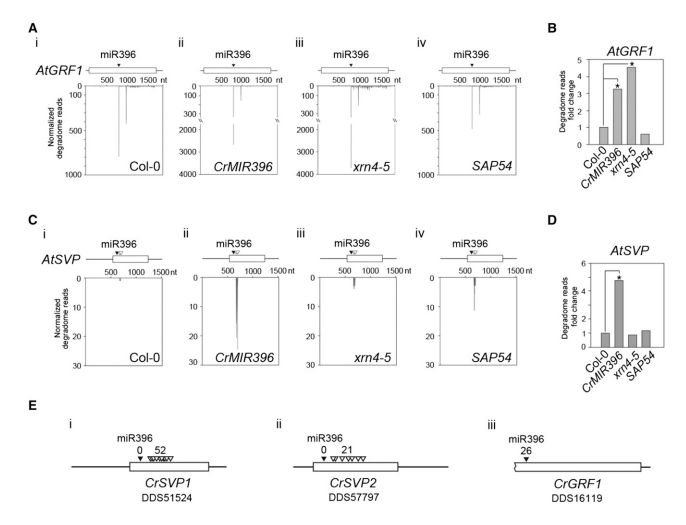


Figure 8. PHYL1 alters the expression of miR396 and its targets. A, AtSVP and AtGRF1 expression levels in Col-0, PHYL1, and svp32 plants were detected by real-time RT-PCR. Error bars represent se (n = 9). Relative expression levels were normalized to the level of AtUBQ. Relative expression levels were significantly different from those of the Col-0 plants in each RNA sample, based on Student's t test: *, P < 0.05; and **, P < 0.01. B, Detection of SVP gene expression in floral bud tissue by in situ hybridization. The buds of Col-0 (i) and PHYL1 (ii) plants were sectioned and hybridized with an antisense probe for SVP detection. The sense probe for AtSVP was used as a negative control (iii and iv). The black arrows indicate the AtSVP signal at the sepal, and the red arrows indicate the AtSVP signals at the basal part of the petal. an, Anther; pe, petal; se, sepal; st, stigma. Bar = 50 μ m.



CrSVP1 and CrSVP2, respectively, suggesting the presence of miR396-mediated CrSVP1/2 decay. In contrast, 26 degradome reads of CrGRF1 were detected in the target position, indicating miR396-mediated cleavage (Fig. 9É, image iii, black arrowhead). In addition, one predicted miR396 target gene (AT1G71350) exhibits mismatch pairing at the 5' portion (Supplemental Fig. S7). No significant miR396-cleaved degradome reads at the target site were observed for the gene. However, AT2G31880 has approximately 5-fold more degradation reads in CrMIR396 plants (Supplemental Fig. S7, B and C). In contrast, under the miR396-decreasing condition, AT2G31880 expression was increased 50% in SAP54 plants, which suggests the presence of miR396-mediated AT2G31880 down-regulation (Supplemental Fig. S7D). In summary, based on the 5' portion pairing differences between miR396 and its targets, miR396-mediated SVP or AT2G31880 down-regulation might differ from the

GRF1 control. In addition, this phenomenon is conserved in various plant species.

DISCUSSION

Multifunctional effectors are common in many pathogens. In addition to MTF degradation, we demonstrated that PHYL1 effector can up-regulate *SVP* MTF expression and alter miRNA expression to switch the phase transition back to the vegetative stage. In addition, we also provided evidence to show that miR396 mediated *SVP* mRNA decay.

The Effectors of PnWB Alter Gene and miRNA Expression in Host Plants

The virescence and phyllody symptoms on phytoplasmainfected plants are remarkable phenotypes. Phytoplasma

infection alters the physiological and morphological characteristics of the host cells to enhance pathogenic multiplication and infection in the *C. roseus* plants (Sugio et al., 2011). To revert the reproductive stage to the vegetative stage, PHYL1/SAP54 effectors suppress positive regulators of flowering but enhance the expression levels of negative regulators, such as *CrSVP1* and *CrSVP2*. However, *CrSOC1.1*, *CrSOC1.2*, and *CrAP1.2*, the orthologs of which function as positive regulators in flowering in Arabidopsis, were not repressed by PnWB. This phenomenon can be explained by two forces that originate from the pathogen and host to counteract each other; they are abnormally coexpressed in the floral meristem, resulting in leafy flower formation (Liu et al., 2014).

Rather than altering gene expression, the miRNA expression levels were also altered during PnWB infection. Interestingly, the miRNA expression patterns in PnWB-infected S4 leafy flowers differ in *PHYL1* and *SAP54* plants. We hypothesize that the other effectors or bacterial proteins in PnWB directly or indirectly control *CrMIR* gene expression, resulting in different miRNA expression patterns between PnWB-infected plants and transgenic plants. Moreover, the PHYL1/SAP54 that triggers MTF degradation might alter abnormal miRNA expression to cause a synergistic effect, thereby resulting in the leafy flower phenotype.

Like SAP54 or PHYL1 of onion yellows phytoplasma mutant plants, PHYL1 of PnWB effector causes leafy flowers in transgenic Arabidopsis plants. Moreover, the miRNA expression patterns of *PHYL1* plants (PHYL1 of PnWB) were also consistent with those of SAP54 plants. SAP54 and PHYL1 of onion yellows phytoplasma plants mediate AtAP1, AtSEP3, and AtCAL proteasomal degradation (MacLean et al., 2014; Maejima et al., 2014); therefore, MTF degradation might result in abnormal miRNA expression. These results indicate that these effectors contain common motif(s) for triggering leafy flower formation and altering miRNA expression. Furthermore, PHYL1 mediates the down-regulation of miR396 expression; however, it did not interfere with miR396 biogenesis in vivo. PHYL1 might trigger MTF degradation or alter the expression levels of other factors that regulate miR396 expression.

In addition, the *atmir396a-1* mutant exhibited *AtSVP* accumulation without manifesting any abnormal flower phenotype. Our explanation for this observation is that the functional redundancy of miR396b might play a role in regulating *AtSVP*. Moreover, the 0.5-fold accumulation of *AtSVP* relative levels in *atmir396a-1* plants might not be sufficient to trigger the abnormal flower phenotype, whereas the 3- and 90-fold accumulation of *AtSVP* in *PHYL1* and *AtSVP* plants, respectively, might trigger this phenotype (Fig. 8; Supplemental Fig. S4B). In contrast, *AtMIR396a* and *CrMIR396* plants did not show the same early-flowering phenotype as the *svp32* mutant, possibly because miR396 also regulates other genes; therefore, the synergistic effect resulted in a milder late flowering.

The 5' Portion of miR396 Determines Target RNA Degradation by miRNA Cleavage or mRNA Decay

The degradome patterns indicated that miR396mediated SVP and GRF1 degradation profiles differ because of the discrepant pairing on the 5' portion of the miRNA. The 5' portion (i.e. the fourth and fifth positions) of the miRNA is critical for miRNAmediated cleavage, whereas the 19th and 20th positions are nonessential (Lin et al., 2009). The 5' mismatch of miRNA reduces the efficiency of target RNA cleavage and may trigger mRNA decay (Lin et al., 2009; Bazzini et al., 2012; Djuranovic et al., 2012), which may explain the failure to confirm the miR396 target site of SVP by 5' RACE (Chorostecki et al., 2012). Indeed, the degradome profiles showed that CrSVP1, CrSVP2, and AtSVP have nonspecific mRNA decay, whereas a clear degradome pick was shown on the target site of CrGRF1 and AtGRF1. In miRNA regulation of mRNA by translation repression in animal models, miRNA-mediated silencing complex bound of mRNA is typically degraded by another endogenous RNase system (Stefani and Slack, 2008; Bazzini et al., 2012; Djuranovic et al., 2012). In plants, the miR398mediated cleavage of the COPPER/ZINC SUPEROXIDE DISMUTASE (CSD1 and CSD2) family, which exhibits a mismatch at the fourth position of miR398, demonstrated that miR398 regulates CSD2 expression through translation repression (Sunkar et al., 2006; Li et al., 2013). Therefore, we hypothesize that SVP mRNA decay might be triggered by miR396-mediated translation inhibition.

PHYL1-Mediated MTF Degradation Might Interfere with the Heterodimer Formation of the SVP Complex

PHYL1/SAP54 interacts with the RAD23 family to trigger MTF (AP1, SEP3, and CAL) degradation,

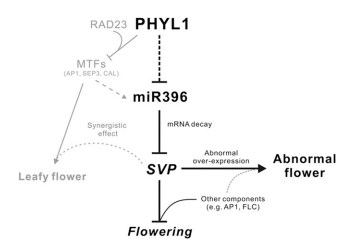


Figure 10. Model for PHYL1-regulated miR396-mediated *SVP* expression in the flowering pathway resulting in the abnormal flower phenotype.

resulting in the formation of leafy flower phenotypes (MacLean et al., 2014; Maejima et al., 2014). However, AtSVP interacts with AP1 or other components to form different complexes that regulate floral gene expression (Gregis et al., 2009). Therefore, the lack of AP1 in PHYL1/SAP54 plants might prevent AtSVP from forming a heterodimer complex, thereby resulting in misregulation during the phase transition and floral organ identity. In addition, the degrees of phyllody and virescence of SAP54 plants in the rad23 family mutant background were reduced, especially in the rad23BCD triple mutant that has no strong leafy flower formation in the presence of phytoplasma infection or SAP54 overexpression (MacLean et al., 2014). Interestingly, rad23BCD mutant plants have shown the abnormal flower (five to six petals) phenotype (MacLean et al., 2014) identical with the AF-I of PHYL1 and AtSVP plants. Therefore, these data implied that the gain of function of the AtSVP-AP1 complex might also affect flower formation, while AP1 shows no degradation in the *rad23BCD* mutant background (Fig. 10). In addition, the SVP complex has species specificity with regard to interference with flowering time and flower development in Arabidopsis, because only AtSVP plants show the late-flowering and AF-I phenotypes in Arabidopsis, whereas other SVP plants have normal flowering time and normal flowers (except AdSVP plants), similar to Col-0 plants.

A Working Model for PHYL1 Repression of miR396-Mediated SVP Regulation

We propose a model for PnWB repression of the miR396-mediated *SVP* silencing pathway (Fig. 10). Multiple functions of PHYL1 trigger positive regulators of MTF (AtAP1, AtSEP3, and AtCAL) proteasomal degradation, resulting in leafy flower formation. PHYL1-mediated MTF degradation also affects many floral homeotic gene expression levels and miRNA expression levels, including the up-regulation of *AtSVP* and the repression of miR396. MiR396 regulates *SVP* through an mRNA decay approach that might be triggered by translation inhibition. Therefore, the down-regulation of *SVP* to change the complex formation to repress flowering and cause an abnormal flower phenotype.

CONCLUSION

This study provides evidence to demonstrate that PHYL1/SAP54 effectors can change miRNA expression levels. In addition, we also demonstrate that miR396 can down-regulate *SVP*, and the overexpression of *AtSVP* causes the abnormal flower phenotype. In the future, we will explore how PHYL1 controls these miRNA gene expression levels and examine symptom development with regard to the pathogen effector-host factor interaction at the posttranscriptional level.

MATERIALS AND METHODS

Plant Materials and Phytoplasma Infection

Arabidopsis (*Arabidopsis thaliana*) seeds were surface sterilized and chilled at 4°C for 2 d prior to sowing on Murashige and Skoog (MS) medium with or without suitable antibiotics for selection. One-week-old seedlings were germinated on MS plates and transferred to soil. *Catharanthus roseus* and *Nicotiana benthamiana* seeds were sown in soil. All seedlings and plants were maintained in a growth chamber or greenhouse (16 h of light/8 h of dark, 20°C–25°C).

Two-month-old *C. roseus* plants were inoculated with PnWB phytoplasma via grafting. The PnWB-infected plants developed virescence symptoms 3 to 4 weeks post grafting, whereas phyllody symptoms developed at 5 to 6 weeks post grafting.

Screening of atmir396a Mutants of Arabidopsis

Mutant lines of atmiR396a-1 (SALK064047) for T-DNA insertion were obtained from The Arabidopsis Information Resource (http://www.arabidopsis.org/). Genomic DNA from 1-week-old seedlings was used for genotyping. The primers LP and RP were used to detect the wild-type genome region of the AtMIR396a gene, whereas the primers LBb1.3 and RP were used to detect the T-DNA insertion region. The primers used for genotyping are listed in Supplemental Table S1.

Gene Construction and Transgenic Plants

The genes AtSVP, CrSVP1, CrSVP2, AtMIR396a, CrMIR396, and PnWB PHYL1 were identified based on the Arabidopsis (The Arabidopsis Information Resource) and C. roseus transcriptome databases (Chang, 2012; Liu et al., 2014). AtSVP was cloned via RT-PCR amplification using the primers F-AtSVP and R-AtSVP. For AtGRF1, the 5' end of AtGRF1 (792 nucleotides) was cloned via RT-PCR amplification using the primers F-AtGRF1 and R-AtGRF1-792. CrSVP1 was cloned via RT-PCR amplification using the primers 51524_F and 51524_R. CrSVP2 was cloned via RT-PCR amplification using the primers 57797_F and 57797_R. AtMIR396a was cloned via RT-PCR amplification using the primers F-AtMIR396a and R-AtMIR396a. CrMIR396 was cloned via RT-PCR amplification using the primers 34004-29-F and 34004-235-R. The PHYL1 gene was identified from the PnWB genome sequence (Chung et al., 2013) and amplified from the cDNA of PnWB-infected C. roseus by RT-PCR with the primers FG-PHYL1-Pn and RG-PHYL1-Pn. PCR fragments were cloned into the pENTR/D-TOPO vector (Invitrogen) according to the manufacturer's instructions to generate pENTR-CrSVP1, pENTR-CrSVP2, pENTR-CrMIR396, and pENTR-PHYL1.

For the miR396 target-site mutation of *AtSVP*, the *AtSVP* cDNA was used as a template for PCR-based mutagenesis with MAtSVPm2-118 and PAtSVPm2-70. The PCR fragments were cloned into pENTR/D-TOPO vector to generate pENTR-AtSVPm.

For the reporter assay, the $AtSVP_{129}$ and $AtSVP_{129m}$ DNA fragments (nucleotides 1–129 of the 5' end of either AtSVP or AtSVPm) were amplified by PCR from pENTR-AtSVP as well as pENTR-AtSVPm with F-AtSVP and MAtSVP-129 primers. In addition, the $GRF1_{129}$ DNA fragment was amplified by PCR from pENTR-AtGRF1 with PAtGRF1-731 and MAtGRF1-817 primers. These PCR fragments were cloned into the pENTR/D-TOPO vector to generate pENTR-AtSVP₁₂₉, pENTR-AtSVP_{129m}, and pENTR-AtGRF1₁₂₉. Gateway system procedures were used to transfer these genes to a pBA-DC-YFP binary vector to generate pBA-AtSVP₁₂₉-YFP, pBA-AtSVP_{129m}-YFP, and pBA-AtGRF1₁₂₉-YFP.

These binary plasmids were transformed into an *Agrobacterium tumefaciens* ABI strain for agroinfiltration or used for transgenic Arabidopsis transformation using a floral dip procedure (Zhang et al., 2006). The transformant lines were screened on MS medium containing $10 \ \mu g \ mL^{-1}$ Basta (Sigma).

The artificial miRNA precursor that targets the *GFP* gene (*preamiR-GFP4*) was constructed based on the miR159 precursor of Arabidopsis via PCR mutagenesis using the primers PamiR-FG and MamiR. The PCR product was cloned into a pENTR/D-TOPO vector to generate pENTR-amiR-GFP4. Gateway system procedures were used to transfer *preamiR-GFP4* to a pBA-DC-HA binary vector (Zhang et al., 2005) and generate pBA-amiR-GFP4. The primers used for cloning are listed in Supplemental Table S1.

Transient Expression via Agroinfiltration

An A. tumefaciens ABI strain-containing binary vector was incubated in Luria-Bertani medium containing 10 mm MES, pH 5.6, 40 μ m acetosyringone,

 $100~\mu g~mL^{-1}$ spectinomycin, and $50~\mu g~mL^{-1}$ kanamycin at $28^{\circ}C$ for 16 h. The A. tumefaciens was suspended, the absorbance was adjusted to optical density at 600~nm=0.5 using an appropriate buffer ($10~mm~MgCl_2$ and $150~\mu m$ acetosyringone), and the A. tumefaciens was incubated at room temperature for 3 h (Lin et al., 2013). Four days after agroinfiltration, the infiltrated leaves of the N. benthamiana plants were collected and analyzed by semi-RT-PCR and northern blot.

For the observation of reporter assay results, YFP fluorescence in the infiltrated leaf was monitored using the Leica TCS SP5 II confocal laser-scanning microscope (Joint Center for Instruments and Research, College of Bioresources and Agriculture, National Taiwan University) equipped with a multiline argon laser with a filter set for YFP fluorescence (excitation filter, Acousto-optic Tunable filter 488; emission bandwidth, 496 to 574 nm; PMT2 offset [-1]/gain [895]). All images were graphically arranged used Adobe Photoshop CS3 software.

Total RNA Extraction and Small RNA Detection

Total RNA was extracted from the plant tissues using TRIzol reagent (Invitrogen) according to the manufacturer's protocol. For small RNA northern blotting, $10~\mu g$ of total RNA was separated on a 15% polyacrylamide/1× Trisborate/EDTA/8 M urea gel and transferred to a Hybond-N $^+$ membrane. The miR396-AS DNA probe (5'-CAGTTCAAGAAAGCTGTGGAA-3') and the amiRGFP4-AS DNA probe (5'-TACTCCAATTGGCGATGGCCC-3') were end labeled with [γ - 32 P]ATP using T4 polynucleotide kinase (New England Biolabs). Free radioisotopes were filtered out using a mini Quick Spin Oligo Column (Roche). The membrane was hybridized with ULTRAhyb-Oligo hybridization buffer (Ambion) at 42° C for 16 h, and the signal was detected using X-film (GE Healthcare) at -80° C for 16 h.

Real-Time RT-PCR, Semi-RT-PCR, and Northern Blot

For real-time RT-PCR, cDNA was first reverse transcribed from total RNA using oligo(dT) $_{20}$ and the SuperScript III First Strand Synthesis System (Invitrogen). Real-time PCR was performed using a LightCycler 480 instrument (Roche) and three sets of primers for AtSVP, AtGRF1, and AtUBQ expression detection. The primers used for real-time RT-PCR are listed in Supplemental Table S2. For semi-RT-PCR detection, the primers F-CrSVP1_A and R-CrSVP1_C were used to detect CrSVP1. The primers F-AtSVP and R-AtSVP were used to detect AtSVP. The primers F-AdSVP2 and R-AdSVP2 were used to detect AtSVP. The primers CrUBQc-F and CrUBQc-R were used to detect CrUBQ. The primers NbActin-F1 and NbActin-R1 were used to detect NbACTIN. The primers AtUBQ-F1 and AtUBQ-R1 were used to detect AtUBQ. The primers used for semi-RT-PCR are listed in Supplemental Table S3.

Statistical Analyses and Deep Sequencing

The AtSVP, AtGRF1, and AtUBQ transcripts were quantified via a relative cycle threshold (Ct) method. All experiments were performed with three independent replicates to compensate for possible loading errors. The relative expression levels were calculated based on the $\Delta\Delta$ Ct value, and each sample was normalized according to the expression level of AtUBQ. The miR396 expression levels were normalized via U6 internal/loading controls, which were quantified using ImageQuant 5.2 (GE Healthcare). The $\Delta\Delta$ Ct values were used to explain the fold changes in the expression levels.

The small RNA profiles of the PHYL1 plant, SAP54 plant, and HF and S4 flowers were analyzed using Illumina HiSeq 2000. The degradome profiles of the Col-0 plant, CrMIR396 plant, xrn4-5 mutant, SAP54 plant, and HF of C. roseus plant were also analyzed using Illumina HiSeq 2000. The miRNA and degradome profiles were analyzed using CLC Genomics Workbench 5.5.2 (CLC Bio) and the ContigViews analysis platform (www.contigviews.bioagri.ntu.edu. tw). The read normalization for the miRNA and degradome assays was calculated by the ratio of total reads across various samples to Col-0 plants. The raw small RNA reads reported here have been submitted to the National Center for Biotechnology Information Short Read Archive under accession numbers SRR1818286 (normal flower of Col-0 plant), SRR1818303 (leafy flower of PHYL1 plant), and SRR1818289 (leafy flower of SAP54 plant). The raw degradome reads reported are available in the Short Read Archive under accession numbers SRR1821048 (flower of Col-0 plant), SRR1821008 (flower of CrMIR396 plant), SRR1821172 (flower of $xm4-\bar{5}$ mutant), SRR1821156 (leafy flower of SAP54plant), and SRR1821217 (HF of C. roseus plant).

In Situ Hybridization

A digoxigenin (DIG)-labeled SVP RNA probe (1,117–1,416 bp) was produced using an in vitro transcription kit (Roche) following the manufacturer's protocol. The embedded Col-0 and PHYL1 inflorescence tissues were sectioned to 8- μ m thickness and hybridized with DIG-labeled RNA probes. The hybridization signal was detected using an anti-DIG antibody conjugated to alkaline phosphatase (Roche) with nitroblue tetrazolium/5-bromo-4-chloro-3-indolyl phosphate as the substrate. The results were examined with a bright-field microscope as purple dotted signals. The protocol followed that of a previous study (Wang et al., 2008).

Supplemental Data

The following supplemental materials are available.

Supplemental Figure S1. Alignment and comparison of the parallel positions of the miR396 target sites from MADS box-containing Arabidopsis genes.

Supplemental Figure S2. In vivo analysis of the miR396-mediated *CrSVP1*, *CrSVP2*, and *AtGRF1* cleavage efficiency by transient expression.

Supplemental Figure S3. In vivo analysis of the miR396-mediated *CrSVP1* expression efficiency by transient expression.

Supplemental Figure S4. Detection of different *SVP* genes in various transgenic plants.

Supplemental Figure S5. Amino acid sequence alignment of AtSVP, CrSVP1, CrSVP2, and AdSVP2.

Supplemental Figure S6. Degradome analysis of Arabidopsis.

Supplemental Figure S7. Degradome data regarding the mismatches in the 5' portion of the miR396 pairwise region.

Supplemental Table S1. Primers used in this study.

Supplemental Table S2. Primers for real-time RT-PCR.

Supplemental Table S3. Primers for semi-RT-PCR.

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