

Regulation of flowering time via miR172-mediated *APETALA2-like* expression in ornamental gloxinia (*Sinningia speciosa*)^{*#}

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Abstract: We investigated the microRNA172 (miR172)-mediated regulatory network for the perception of changes in external and endogenous signals to identify a universally applicable floral regulation system in ornamental plants, manipulation of which could be economically beneficial. Transgenic gloxinia plants, in which miR172 was either overexpressed or suppressed, were generated using *Agrobacterium*-mediated transformation. They were used to study the effect of altering the expression of this miRNA on time of flowering and to identify its mRNA target. Early or late flowering was observed in transgenic plants in which miR172 was overexpressed or suppressed, respectively. A full-length complementary DNA (cDNA) of gloxinia (*Sinningia speciosa*) *APETALA2-like* (*SsAP2-like*) was identified as a target of miR172. The altered expression levels of miR172 caused up- or down-regulation of *SsAP2-like* during flower development, which affected the time of flowering. Quantitative real-time reverse transcription PCR analysis of different gloxinia tissues revealed that the accumulation of *SsAP2-like* was negatively correlated with the expression of miR172a, whereas the expression pattern of miR172a was negatively correlated with that of miR156a. Our results suggest that transgenic manipulation of miR172 could be used as a universal strategy for regulating time of flowering in ornamental plants.

Key words: Flowering time; Transgenic gloxinia; MicroRNA172; *APETALA2-like* (*AP2-like*)
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1 Introduction

Plants rely on a complex genetic regulatory network for the perception of changes in external and endogenous signals to select the appropriate time of flowering. Within this vast regulatory network, microRNAs (miRNAs) play pivotal roles by controlling the expression of key flowering genes.

All miRNAs are non-coding RNAs of 20–24 nucleotides long that specifically regulate the expression of target genes at the transcriptional or translational level in eukaryotes. Multiple developmental events are regulated by miRNAs in plants (Chen, 2005). Among the numerous miRNA families, only three, namely miR156, miR172, and miR159/miR319, are associated with the regulation of flowering time (Terzi and Simpson, 2008). The overexpression of miR159/miR319 delays flowering: miR159 is mainly involved in gibberellin signaling pathways (Achard et al., 2004; Terzi and Simpson, 2008; Li et al., 2013).

In *Arabidopsis*, there are eight loci that encode the members of the miR156 family, including

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miR156a–miR156j (Morea et al., 2016). The expression level of miR156 is highest in plant embryos and plantlets, and it gradually decreases during plant growth (Hong and Jackson, 2015). The main target genes of the miR156 family are the SQUAMOSA PROMOTER BINDING PROTEIN (SBP)-box transcription factors (Ferreira et al., 2014). In *Arabidopsis*, miR156 regulates the timing of the transition from the juvenile to the adult stage by repressing the transcript levels of SQUAMOSA PROMOTER-BINDING PROTEIN-LIKE (SPL) transcription factors (Yang et al., 2011). *SPL9*, a target of miR156, binds directly to the regulatory region of miR172b to promote its expression and induce flowering (Wu et al., 2009). miR172, which is downstream of miR156, promotes adult epidermal identity. Both miR156 and miR172 are associated with aging in plants.

MiR172 has been predicted to suppress a set of transcription factors, including APETALA2 (AP2) and related proteins, to promote flowering (Zhu and Helliwell, 2011). In *Arabidopsis*, the overexpression of miR172 caused early flowering and floral defects under long-day conditions. An early-flowering mutant, *early activation tagged dominant (eat-D)*, exhibited a phenotype similar to that of the miR172 overexpression line (Aukerman and Sakai, 2003). The overexpression of *TARGET OF EAT1 (TOE1)*, which is a target gene of miR172, resulted in delayed flowering onset. Mutant plants deficient in *TOE1* flowered slightly earlier, but the effect was not significant compared with the wild type (Aukerman and Sakai, 2003). Although flowering occurred significantly earlier in *toe1 toe2* double mutants, the extent of early flowering was less significant than in miR172-overexpressing plant lines (Aukerman and Sakai, 2003).

Recent studies have shown that the overexpression of *SCHLAFMUTZE (SMZ)* and *SCHNARCHZAPFEN (SNZ)* leads to delayed flowering, whereas *smz snz* double mutants do not show significant changes in flowering time compared with the wild type (Mathieu et al., 2009). *Toe1 toe2 smz snz* quadruple mutants flower significantly earlier than *toe1 toe2* double mutants, but the extent of early flowering was less significant than in the miR172-overexpressing plant lines (Yamaguchi and Abe, 2012). This indicates that the *AP2-like* gene family has a certain degree of functional redundancy in the regulation of flowering

time (Mathieu et al., 2009). In loss-of-function sextuple mutants in which all *AP2-like* genes were affected, flowering occurred earlier than in the miR172-overexpressing transgenic plant line, indicating that all *AP2-like* genes are flowering inhibitors (Mathieu et al., 2009; Yant et al., 2010).

Gloxinia (*Sinningia speciosa*) is an herbaceous commercial plant native to south-eastern Brazil, with large, bell-shaped flowers. Because of their long flowering time and colorful flowers, these plants are popular as houseplants. However, the mechanism regulating flowering time is not known. We investigated whether genetic modification of the *miR172* gene can provide a universal way to regulate flowering time in ornamental plants by generating transgenic gloxinia plants in which *miR172* was overexpressed or suppressed. Candidate target genes of miR172 were identified, and a full-length complementary DNA (cDNA) of *AP2-like* was cloned to verify whether it is a target of miR172. We investigated whether the altered levels of miR172 modulated the expression of *AP2-like* genes, and whether the changes would affect the time of flowering under short-day conditions.

2 Materials and methods

2.1 Plant growth conditions

Young leaves cut from cultured plantlets of gloxinia were placed on Murashige and Skoog's (MS) culture medium (pH 5.8) supplemented with hormones at different concentrations (Table S1). The cultures were placed in a chamber under short-day conditions (8-h light/16-h darkness, photon flux density of 25 $\mu\text{mol}/(\text{m}^2\cdot\text{s})$) at 24 °C for differentiation. For regeneration, differentiated plantlets at the same developmental stage were transferred to root-inducing medium (Table S1). After one month, all the plantlets were transferred to a perlite and vermiculite matrix mixture. To assay the roles of miR172a in regulating flowering time, wild-type and transgenic gloxinia plants were grown under short-day conditions (8-h light/16-h darkness, (24 \pm 1) °C).

2.2 Plasmid construction

Sequences available in GenBank (<http://www.ncbi.nlm.nih.gov/genbank>) suggested that the core

regions of the miR172 family are highly conserved in plants (Fig. 1). Because no information on the SsmiR172a precursor was available in public databases, we employed the AtmiR172a precursor (*Columbia-0*, *Wassilewskija* (*Ws*) ecotype) to generate mature miR172a which we overexpressed in gloxinia under the regulation of cauliflower mosaic virus 35S (CaMV 35S) promoter. To better understand the roles of miR172a in gloxinia, we used target mimicry (35S:*MIMI172*) to suppress the activity of miR172a under the control of *INDUCED BY PHOSPHATE STARVATION1* (*IPS1*), as described in previous reports (Franco-Zorrilla et al., 2007; Li et al., 2013). On the basis of sequences available in the National Center for Biotechnology Information (NCBI) database (<https://www.ncbi.nlm.nih.gov>), primers were designed for the amplification of miR172a precursor from *Arabidopsis* (*Columbia-0*, *Ws* ecotype) cDNA. The sense (5'-TCCCCCGGGTTCTTCTGGTTTGGAGATGGTTAGG-3') and anti-sense primers (5'-ACGCGTCGACCTTGATAAAGACTGCCAAAACC TG-3') contained *Sma*I and *Sal*I sites at their 5'-ends, respectively. The artificial *IPS1* fragment was cloned using forward (5'-CTTACGTCGTAGTAGGAGTTCTAAGATTTCTAGAGGGAGATAA-3') and reverse (5'-AATCTTAGAACTCCTACTACGACGTAAGCTTCGGTTCCCCTCG-3') primers designed for its amplification. Polymerase chain reactions (PCRs)

were carried out using 2× Taq Master Mix (TaKaRa, Dalian, China) following the protocol provided by the supplier. For each PCR, 5 μL of the template was used in a 50-μL reaction mixture containing 25 μL Taq Master Mix, 200 nmol/L of each primer, and 18 μL of triple-distilled water. PCR was conducted for three biological replications of each sample using the following thermal cycles: initial denaturation at 94 °C for 5 min, followed by 30 cycles, each cycle consisting of 94 °C for 15 s, 50–60 °C for 30 s, and 72 °C for 15 s, and final extension at 72 °C for 10 min.

All fragments were cloned into the pCAMBIA13011 plasmid, which contained a CaMV 35S promoter and the terminator of nopaline synthase (Nos), as well as kanamycin- and hygromycin-resistance genes. All recombinant plasmids were transferred into *Agrobacterium tumefaciens* strain EHA105.

2.3 Screening of transgenic gloxinia plants

Transgenic gloxinia plants were generated by *Agrobacterium*-mediated transformation as described previously (Zhang et al., 2008; Li et al., 2013). After four cycles of fortnightly selection (Table S1), hygromycin-resistant plantlets were transferred to root-induction medium without hygromycin. PCR was used to screen the transgenic plantlets with gene-specific primers and genomic DNA.

2.4 Genomic DNA isolation

Genomic DNA was isolated from young leaves of transformed plants using a genomic DNA isolation kit (TaKaRa, Dalian, China), following the manufacturer's protocol, and used for PCR-based screening of the hygromycin-resistance gene.

2.5 Total RNA isolation

Total RNA was extracted from leaves and flower buds using TRIzol reagent (TaKaRa, Dalian, China) according to the manufacturer's instructions. The concentration and purity of the extracted RNA were measured using a BioSpectrometer (Eppendorf, Hamburg, Germany). Reverse transcription was carried out with 1 μg RNA following the instructions provided with the PrimeScript 1st Strand cDNA Synthesis Kit (TaKaRa, Dalian, China). Mature miRNAs were detected using stem-loop reverse transcription PCR (RT-PCR) (Varkonyi-Gasic and Hellens, 2010; Li et al., 2013).

At-miR172a	AGAAUCUUGAUGAUGCUGCAU
Sl-miR172a	AGAAUCUUGAUGAUGCUGCAU
Al-miR172a	AGAAUCUUGAUGAUGCUGCAU
Aq-miR172a	AGAAUCUUGAUGAUGCUGCAU
Bo-miR172a	AGAAUCUUGAUGAUGCUGCAU
Br-miR172a	AGAAUCUUGAUGAUGCUGCAU
Eg-miR172a	AGAAUCUUGAUGAUGCUGCAU
Gm-miR172a	AGAAUCUUGAUGAUGCUGCAU
Os-miR172a	AGAAUCUUGAUGAUGCUGCAU
Pt-miR172a	AGAAUCUUGAUGAUGCUGCAU
Vv-miR172a	UGAAUCUUGAUGAUGCUACAU

Fig. 1 Alignment of mature miR172a sequences from different plant species

Sequence alignment of mature miR172a from different species: *Arabidopsis thaliana* (At), *Solanum lycopersicum* (Sl), *Arabidopsis lyrata* (Al), *Aquilegia coerulea* (Aq), *Brassica oleracea* (Bo), *Brassica campestris* (Br), *Elaeis guineensis* (Eg), *Glycine max* (Gm), *Oryza sativa* (Os), *Populus trichocarpa* (Pt), and *Vitis vinifera* (Vv)

2.6 Cloning of full-length gloxinia *APETALA2-like* (*SsAP2-like*) cDNA and sequence analysis

Degenerate primers (forward primer: 5'-TAG GTGGATTTGAYACNGCWCATGC-3' and reverse primer: 5'-CCYAAATAAACRTACTKTTTGCCT-3') were designed on the basis of the conserved region in the *AP2* sequences of *Arabidopsis*, tomato, and grape. The PolyA Tract mRNA Isolation System III (Promega, Madison, WI, USA) was used to purify gloxinia mRNA. On the basis of the partial sequence of the cDNA that was amplified, 5'-rapid amplification of cDNA ends (5'-RACE) and 3'-RACE were performed using the SMARTer RACE cDNA Amplification Kit (Clontech, Mountain View, CA, USA) to obtain the full-length cDNA sequences (5'-RACE primer: 5'-CC ACTCCTCTGAATTTAATGGCTGCC-3'; 3'-RACE primer: 5'-GCCCATCTAGCTTCCCATCTACCAC-3'). These sequences were aligned using Omega 2.0 (Kramer, 2001), according to the instructions available for the software, to obtain the full-length sequence of *SsAP2-like*.

2.7 RNA ligase-mediated 5'-RACE

To verify whether *SsAP2-like* is a direct target of miR172a, modified RNA ligase-mediated (RLM) 5'-RACE was conducted to ascertain the cleavage site of miR172 (Li et al., 2013). A round of 5'-RACE was carried out with a special primer (anti-sense primer: 5'-TCATGAGGGTCTCATGAGAGAGTGGATC-3'). The PCR products were cloned into the T-simple vector and ten independent inserts were sequenced.

3 Results

3.1 Generation of transgenic gloxinia plantlets in which miR172a is overexpressed or suppressed

Distinct bands of predicted size were detected in 24 plantlets of AtmiR172a overexpression lines (*35S:miR172a*) and in 21 transgenic plantlets overexpressing *MIMI72* (*35S:MIMI72*), but no bands were detected in wild-type plants (Fig. S1).

3.2 Influence of altered expression levels of miR172a on the flowering time in gloxinia

Under identical cultivation conditions, the lines overexpressing miR172a exhibited early flowering, whereas the *35S:MIMI72* lines showed a delay in flowering compared with wild-type gloxinia plants

(Fig. 2a). The *35S:miR172a* transgenic lines flowered at Day 73.20 ± 2.16 after transplantation; i.e., flowering occurred approximately 50 d earlier than in wild-type plants, which flowered at Day 120.20 ± 3.15 after transplantation. In contrast, the *35S:MIMI72* lines flowered at Day 126.60 ± 4.28 , which was much later than that of the *35S:miR172a* lines (Fig. 2c). Phenotypic analysis revealed that the *35S:MIMI72* transgenic plants were shorter than wild-type and *35S:miR172a* plants. The leaf angles in the *35S:miR172a* plants were larger than those in the wild-type and *35S:MIMI72* plants (Fig. 2b). *35S:miR172a* plants had fewer leaves than wild-type plants (4.91 ± 0.58 and 7.60 ± 0.50 , respectively; Fig. 2d). While the overexpression of the *AP2* gene family resulted in short plant height and slow growth in *Arabidopsis* (Todesco et al., 2010), *35S:MIMI72* plants had fewer leaves than wild-type plants (6.38 ± 0.50 and 7.60 ± 0.50 , respectively) at flowering. These results indicated that alteration of the expression level of miR172a in gloxinia significantly influenced flowering time.

3.3 Isolation and sequence analysis of *SsAP2-like*

The complex functions of *AP2* prompted us to investigate the possible regulatory roles of miR172 and *AP2* in gloxinia during flower development. Sequence analysis showed that the full-length coding sequence of *SsAP2-like* was 1476 base pairs (bp) (Data S1). Phylogenetic analysis showed that the sequence of *SsAP2-like* was similar to that of *Sesamum indicum* (Fig. 3a). Multiple sequence alignment with other *AP2* sequences demonstrated that a putative complementary site of miR172a existed in the region of 1383 to 1403 nucleotides, in which there were only three mismatched nucleotides (Fig. 3b).

A single band of expected size was amplified by nested PCR using cDNA from flower buds as a template (Fig. 3b). The amplified fragment was inserted into the T-simple vector and ten independent clones were sequenced. The results confirmed that *SsAP2-like* is a target of miR172a in gloxinia, as nine of the ten cloned fragments were cleaved at the position of the 10th nucleotide (Fig. 3b).

3.4 Expression patterns of miR156a, miR172a, and *SsAP2-like* in gloxinia

To investigate the expression patterns of miR156 and miR172 in gloxinia, we assessed the accumulation

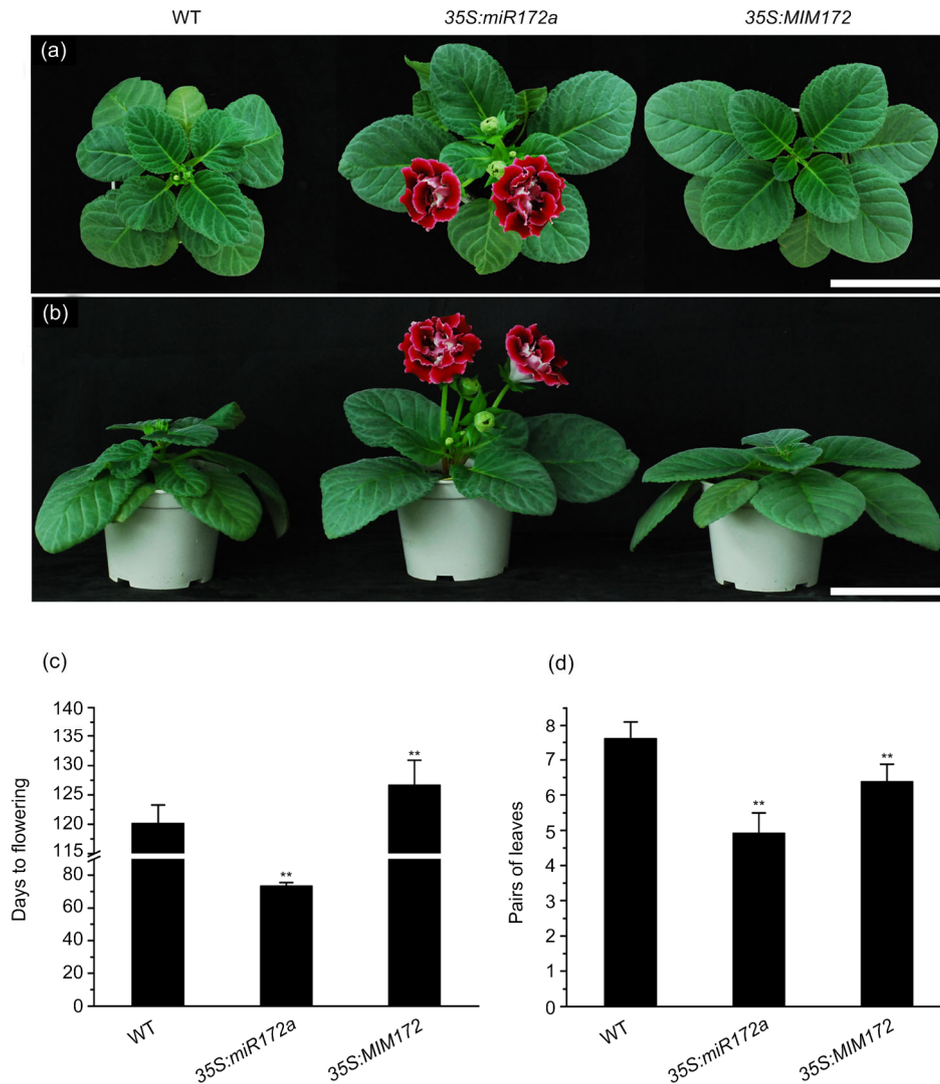


Fig. 2 Influence of flowering time on transgenic gloxinia plants under short-day conditions

(a, b) Photographs were taken at 127 d after transplantation to a mixed matrix. Top views (a) and side views (b) of wild-type (WT), *35S:miR172a*, and *35S:MIM172* plants. Scale bar represents 10 cm. (c) Days to flowering after transplantation. (d) Pairs of leaves at the squaring stage. Data represent the mean±standard deviation (SD) ($n>20$). Double asterisks indicate a significant difference (ANOVA, $P<0.01$) between the wild-type and transgenic plants

of mature miR156a, miR172a, and *SsAP2-like* mRNAs in various tissues. The results showed that miR156a was highly expressed in the leaves (leaf 1), stamens, and carpels of the plantlets, as in the case of *Arabidopsis*. In contrast, the accumulation of miR156a in old leaves (leaves 2 and 3) and sepals was relatively less; in particular, it was nearly absent in the petals (Fig. 4a).

The expression of miR172a was negatively correlated with that of miR156a. With the growth of the plants, the expression level of miR172a in the leaves increased, with the highest accumulation being

observed in the squaring-stage leaves (leaf 3). However, miR172a accumulated relatively less in sepals, petals, and stamens; it was nearly absent in the carpels (Fig. 4b).

SsAP2-like was highly expressed in leaf 1, sepals, petals, and carpels of the plantlet, whereas in the old leaves (leaves 2 and 3) and stamens, the accumulation of *SsAP2-like* was relatively less (Fig. 4c). The expression pattern of *SsAP2-like* was in accordance with that of miR156a, but was negatively correlated with the expression of miR172a in gloxinia.

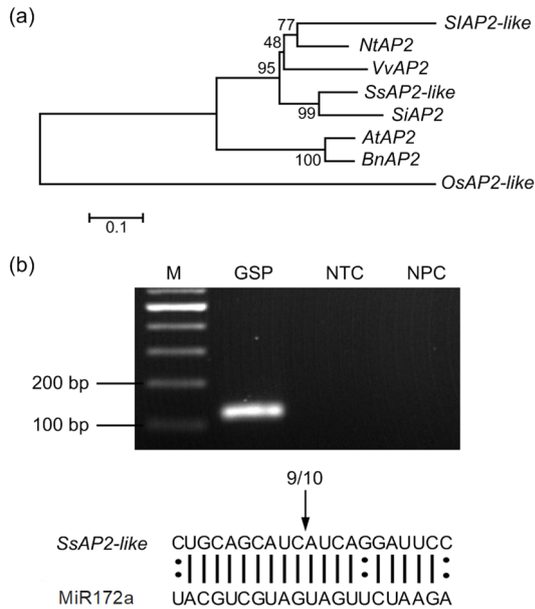


Fig. 3 Analysis of *SsAP2-like* nucleotide sequence and confirmation of miR172a-guided cleavage site in *SsAP2-like* mRNA

(a) Phylogenetic analysis of *SsAP2-like* nucleotide sequences from *Solanum lycopersicum* (Sl), *Nicotiana tabacum* (Nt), *Vitis vinifera* (Vv), *Sesamum indicum* (Si), *Arabidopsis thaliana* (At), *Brassica napus* (Bn), and *Oryza sativa* (Os). Phylogenetic tree was generated by the neighbor-joining method and the *p*-distance model using MEGA4 software. The numbers represent the bootstrap values. (b) Experimental confirmation of miR172 cleavage site in *SsAP2-like* mRNA. The cleavage site was determined by RLM 5'-RACE. M: marker; GSP: gene-specific primer; NTC: no-template control; NPC: no primer control. The arrow indicates the cleavage site of *SsAP2-like* and the number above the sequences indicates the ratio of 5'-RACE clones

3.5 Effects of overexpression and suppression of miR172a on *SsAP2-like* expression in transgenic gloxinia plants

Compared with the wild type, the relative expression levels of mature miR172a in leaves and flower buds were dramatically increased in the 35S:*miR172a* lines and decreased in the 35S:*MIM172* lines (Fig. 5a). These results revealed that we had successfully generated transgenic gloxinia plants with a substantial alteration in the miR172a level in various tissues.

The expression levels of *SsAP2-like* in the 35S:*miR172a* lines were significantly decreased, whereas the levels of *SsAP2-like* were substantially increased in the 35S:*MIM172* lines; this was con-

sistent with the expression pattern of *SsAP2-like*, which was negatively regulated by miR172a (Fig. 5b). These results indicate that the accumulation of *SsAP2-like* mRNA is associated with decreasing expression of miR172a in transgenic gloxinia plants, suggesting that the expression level of *SsAP2-like* is negatively regulated by miR172 in gloxinia.

4 Discussion

The mechanisms of flowering transition in model plants have been extensively investigated and the results have provided a theoretical foundation for research based on the overexpression or inhibition of specific key genes (Zhang et al., 2008). An increasing number of miRNAs have been discovered in plants, and their effects on development and response to environmental change have been studied (Huijser and Schmid, 2011; Teotia and Tang, 2015). However, reports on modifying specific target genes to control flowering time in a species with unknown genomic information are relatively rare and there is a need for more research on the mechanisms of floral transition in economically important ornamental plants. We altered the expression levels of miR172 to affect the accumulation of its target genes, while not manipulating a single gene associated with flowering, and were able to identify an effective pathway for the regulation of flowering time in a species with unknown genomic background.

MiR172 is highly conserved and controls plant growth and development by regulating the expression level of *AP2-like* mRNA, or by inhibiting the translation of *AP2-like* genes (Aukerman and Sakai, 2003; Zhu et al., 2009). Multiple reports have revealed that the overexpression of miR172 causes early flowering under long-day conditions (Aukerman and Sakai, 2003; Lee et al., 2014). The role of miR172 on flowering time regulation prompted us to explore its function in gloxinia during flower development under short-day conditions. In most commercial plants, genomic information is still limited, so we employed the *AtmiR172* precursor to generate transgenic lines that overexpress miR172a. We used the mimicry target approach to obtain miR172-suppressed transgenic lines in which endogenous miR172 activity was eliminated. We identified a member of the *SsAP2-like*

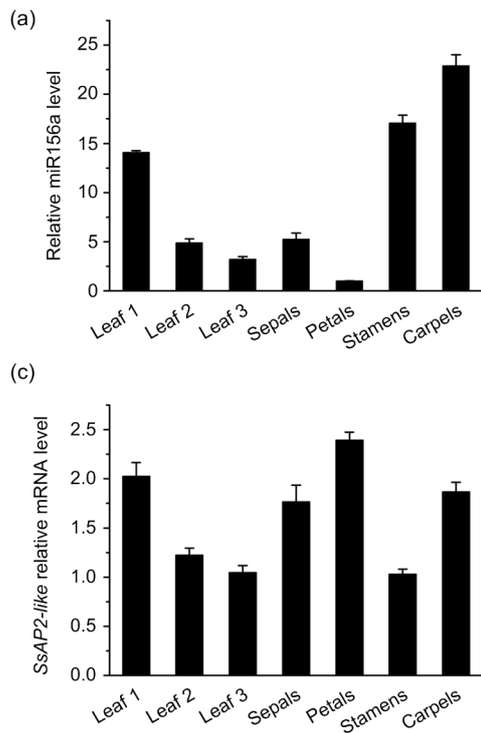


Fig. 4 Analysis of the expression patterns of mature SsmiR156a, SsmiR172a, and *SsAP2-like* in various tissues by RT-qPCR

(a) Accumulation of mature SsmiR156a in various tissues of gloxinia plants. (b) Accumulation of mature SsmiR172a in different tissues of gloxinia plants. (c) Relative mRNA levels of *SsAP2-like* in various tissues of gloxinia plants. Leaf 1: juvenile-phase leaf; Leaf 2: adult-phase leaf; Leaf 3: squaring-stage leaf. Mean values were obtained from three independent samples. Error bars represent SD

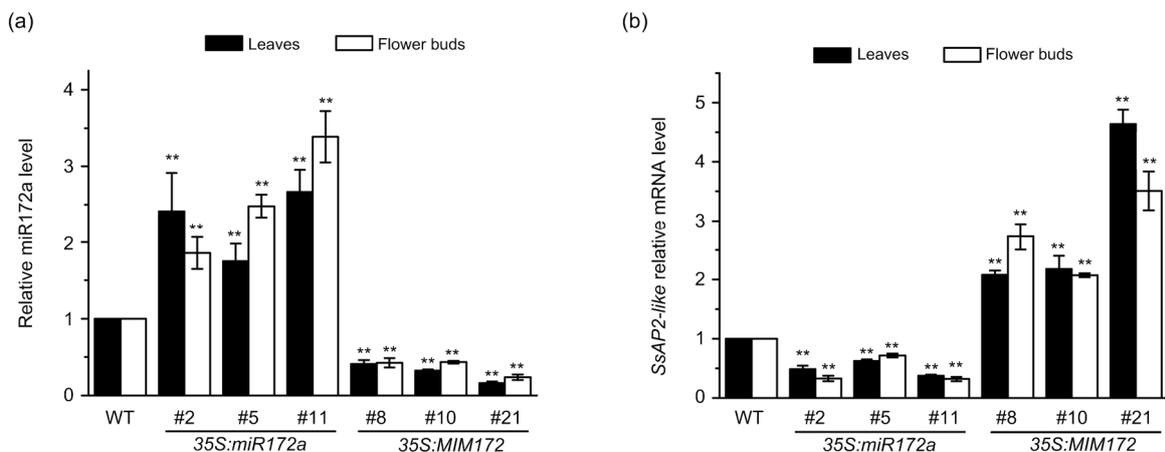


Fig. 5 Relative expression levels of mature miR172a and *SsAP2-like* in wild-type (WT) and transgenic gloxinia plants (a) Transcript levels of mature miR172a in adult-stage leaves and flower buds. (b) Expression levels of *SsAP2-like* in adult-stage leaves and flower buds. Mean values were obtained from three independent samples. Error bars represent the SD. Double asterisks indicate significant differences (ANOVA, $P < 0.01$) between the transgenic lines and WT plants

family in gloxinia that contains a sequence complementary to miR172. *AP2* gene family sequences are relatively conserved in numerous species. In plants, they are under the regulation of miR172, and are widespread, being involved in a variety of physiological and biochemical processes, such as plant growth, floral organ development, and signal transduction. Based on sequence alignment and phylogenetic

analysis, we found that *SsAP2-like* and *SiAP2* had the closest phylogenetic relationship. The *SsAP2-like* mRNA expression levels showed a negative relationship with the abundance of miR172 in both vegetative and floral tissues, consistent with previous findings in *Arabidopsis* and soybean (Lee et al., 2014; Wang et al., 2014). We found that alteration of the expression levels of miR172 significantly influenced

flowering time. The overexpression of miR172 accelerated flowering by approximately 50 d, whereas its suppression delayed flowering by approximately six days compared with wild-type gloxinia under short-day conditions. Alteration of the expression levels of miR172 in gloxinia caused drastic changes in the *AP2* mRNA levels. These findings reveal that miR172 regulates flowering time via the suppression of the *AP2* gene family. In previous studies, alterations in the expression levels of miR172 and *AP2* determined floral patterns and fruit defects (Mlotshwa et al., 2006; Yao et al., 2016). However, in the present study, overexpression of miR172 did not cause any change in the flowers.

MiR156 plays a dominant role in the juvenile phase of plant and flower development because it directly promotes the expression of flowering pathway assembly factors and meristem tissue regulators that control flowering time (Xu et al., 2016). MiR156 can control miR172 expression by regulating SBP-box genes, thereby regulating the activity of *AP2-like* genes in inhibiting flowering. In gloxinia, miR156 and miR172 are expressed alternately. In the juvenile phase, miR156 is expressed at higher levels than miR172. As the plant grows, miR156 expression gradually decreases and miR172 expression gradually increases, which is accompanied by a gradual decrease in *AP2* expression. It has been reported that *AP2* has different functions in floral morphological transitions and plant development (Yant et al., 2010). Analysis of the expression profiles of miR156, miR172, and *AP2* in gloxinia showed that the expression of miR156 was consistent with that of *AP2-like* genes, but opposite to that of miR172. This is consistent with the expression profiles of miR156, miR172, and *AP2* in *Arabidopsis*, indicating that the mutually influential relationship between miR156 and miR172 is relatively conserved in the evolution of plants.

5 Conclusions

The manipulation of miR172 expression in gloxinia alters flowering time. Our study provides a new genetic approach for regulating time of flowering by altering the expression levels of miR172 in ornamental plants.

Contributors

Xiao-yan LI performed the experimental research and data analysis, wrote and edited the manuscript. Fu GUO and Sheng-yun MA collected the data. Mu-yuan ZHU, Wei-huai PAN, and Hong-wu BIAN contributed to the study design, data analysis, and editing of the manuscript. All authors read and approved the final manuscript and, therefore, had full access to all the data in the study and take responsibility for the integrity and security of the data.

Compliance with ethics guidelines

Xiao-yan LI, Fu GUO, Sheng-yun MA, Mu-yuan ZHU, Wei-huai PAN, and Hong-wu BIAN declare that they have no conflict of interest.

This article does not contain any studies with human or animal subjects performed by any of the authors.

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List of electronic supplementary materials

- Table S1 Culture medium used in this study
 Table S2 Primers used in the experiments
 Fig. S1 PCR analysis of *Pre-AtmiR172a* and *MIMI172* (35S: *MIMI172*) in transgenic gloxinias
 Data S1 Nucleotide sequence of *SsAP2-like*

中文概要

题目: MiR172 介导的 AP2-like 转录因子表达对大岩桐花期调控的研究

目的: 本文通过研究 microRNA172a (miR172a) 对成花途径中关键靶基因 APETALA2 类 (AP2-like) 的调控作用及其机理分析, 探索通过操纵 miRNA 表达改变观赏植物花期的基因工程分子育种新策略。

创新点: 本研究基于植物 miR172 序列和功能的高度保守性, 通过转基因的方法操纵大岩桐 miR172 的表达, 进而影响 AP2-like 基因的表达, 并起到调控花期的作用。

方法: 本研究借用拟南芥 miR172a 的已知序列构建组成型过表达载体 35S:miR172a 和抑制 miR172 表达的 35S:MIMI172 载体。利用农杆菌介导法成功获得了 35S:miR172a 过表达株系以及抑制 miR172 作用的 35S:MIMI172 株系, 并利用 cDNA 末端快

速扩增技术 (rapid amplification of cDNA ends, RACE) 克隆得到了大岩桐 *AP2-like* cDNA 全序列, 并通过实时荧光定量聚合酶链式反应技术 (qPCR) 检测转基因株系中 *AP2-like* 的表达变化。

结论: 短日照条件下, *35S:miR172a* 过表达株系花期比野生型提前 (47.00 ± 2.16) 天; *35S:MIM172* 株系花期延迟 (7.00 ± 4.28) 天。在 *35S:miR172a* 过表达株系中 miR172 的表达水平明显上升, 其靶基

因 *SsAP2-like* 的表达量明显下降; *35S:MIM172* 株系中 miR172 的积累水平受到抑制, 而 *SsAP2-like* 的表达量明显上升, 表明 miR172 介导调控 *SsAP2* 的表达对大岩桐成花转变具有促进作用。通过改变 miR172 的表达调控关键靶基因进而改变花期的方法可以作为调节观赏植物开花时间的有效策略。

关键词: 花期; 大岩桐; MicroRNA172; *APETALA2-like* (*AP2-like*)