



# Article The Artificial Promoter *rMdAG2I* Confers Flower-specific Activity in *Malus*

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**Abstract:** Genetic modifications of floral organs are important in the breeding of *Malus* species. Flower-specific promoters can be used to improve floral organs specifically, without affecting vegetative organs, and therefore developing such promoters is highly desirable. Here, we characterized two paralogs of the *Arabidopsis thaliana* gene *AGAMOUS* (*AG*) from *Malus domestica* (apple): *MdAG1* and *MdAG2*. We then isolated the second-intron sequences for both genes, and created four artificial promoters by fusing each intron sequence to a minimal 35S promoter sequence in both the forward and reverse directions. When transferred into tobacco (*Nicotiana benthamiana*) by *Agrobacterium tumefaciens*-mediated stable transformation, one promoter, *rMdAG2I*, exhibited activity specifically in flowers, whereas the other three also showed detectable activity in vegetative organs. A test of the four promoters' activities in the ornamental species *Malus micromalus* by *Agrobacterium*-mediated transient transformation showed that, as in tobacco, only *rMdAG2I* exhibited a flower-specific expression pattern. Through particle bombardment transformation, we demonstrated that *rMdAG2I* also had flower-specific activity in the apple cultivar 'Golden Delicious'. The flower-specific promoter *rMdAG2I*, derived from *M. domestica*, thus has great potential for use in improving the floral characteristics of ornamental plants, especially the *Malus* species.

Keywords: flower-specific promoter; Malus; AGAMOUS; intron

# 1. Introduction

The genus *Malus* comprises more than 30 species with both ornamental and edible uses [1]. Flowering crabapples belonging to this genus are widely used in landscape gardening, due to their varied and attractive flower color, size and fragrance [2]. *Malus* plants are always self-sterile, requiring cross-pollination by insects, such as bees, which is important for fruit production in domesticated apples [3–7]. Flowers are key reproductive organs, because they play important roles in attracting pollinating insects. Therefore, *Malus* flowers are important both for the plants' value as ornamentals and for pollination purposes [8–11]. Improvement of flower traits is a crucial target in breeding *Malus* species.

Transgenic technology has been used in the genetic improvement of a series of flower traits, including flower color, size and stress resistance [12–14]. However, constitutive expression of target genes may have undesirable effects on the vegetative organs. For instance, although cytokinins can be used to regulate flower size, constitutive regulation of cytokinins would also affect vegetative growth [15,16]. A flower-specific promoter could be of great utility by enabling the application of transgenic methods to improve flower traits, specifically without affecting vegetative growth.

In *Arabidopsis thaliana*, the gene *AGAMOUS* (*AG*), initially expressed in the third stage of flower development, is the key factor determining the identity of carpel and stamen [17,18]. The second intron of *AGAMOUS* contains *cis*-regulatory elements influencing the flower-specific expression of target genes [19,20]. In woody plants, Li et al. cloned the second intron of one of the poplar *AG* paralogs, *PTAG2*, and demonstrated that, when fused with the minimal 35S promoter, the reverse version of the *PTAG2* second intron exhibits flower-specific activity [21]. However, a flower-specific promoter derived from the *Arabidopsis AG* second intron does not retain its specificity in tobacco plants [21]. Therefore, isolating the second intron from *Malus* paralogs of *AG* as a tool for the transgenic improvement of flowers in the *Malus* species is needed.

In *M. domestica*, two *MADS*-box genes, *MdMADS15* and *MdMADS22*, have been reported as paralogs of *AGAMOUS* [22]. Furthermore, RNA interference (RNAi) experiments indicate that RNA interference against two apple *AG* paralogs cause male and female sterility, suggesting these genes play important roles in both male and female floral structures [23], similar to the *AG* genes in *Arabidopsis* [24–26]. In this study, we cloned the second introns of two *AG* paralogs, and characterized the promoter activity by fusing them to a  $\beta$ -glucuronidase gene (*uidA*, or *GUS*) in both forward and reverse directions. We reported that the reverse version of the second intron of *MdMADS22* drove the flower-specific expression of target genes both in tobacco and in two *Malus* species, *Malus domestica* (apple) and the ornamental *Malus micromalus* (midget crabapple). This artificial promoter could thus serve as a tool for the genetic manipulation of *Malus* species and other flowering plants.

#### 2. Results and Discussion

#### 2.1. Isolation and Expression Analysis of MdAG1 and MdAG2

To confirm the apple *AGAMOUS* (*AG*) paralogs, we performed a BLAST analysis of the protein sequence of *Arabidopsis AG* against the protein database of the *M. domestica* variety 'Golden Delicious'. We identified two *AG* paralogs, which we named *MdAG1* and *MdAG2*. *MdAG1* and *MdAG2* were reported previously as *MdMADS15* and *MdMADS22*, respectively [22,23]. A phylogenetic tree constructed based on the protein sequences of various well-studied *AG* genes revealed that *MdAG1* and *MdAG2* were (Lustered closely, compared to other *AG* genes (Figure 1).

Next, we characterized the expression patterns of *MdAG1* and *MdAG2* in *M. domestica*. Both *MdAG1* and *MdAG2* exhibited similar floral-organ-predominant expression patterns (Figure 2). Both genes had relatively high expression in pistils and stamens, whereas in vegetative organs such as root, stem and leaf, their expression levels were rarely detectable. In many *Malus* species, fruits develop mainly from receptacles [27]. Interestingly, although both *MdAG1* and *MdAG2* were expressed in receptacles, we did not detect their expression in fruits, indicating that these genes may regulate fruit development in a dynamic fashion [28].



**Figure 1.** Phylogenetic analysis of *AG* protein sequences. The phylogenetic tree was constructed using the neighbor-joining method of MEGA 6.0 software based on the protein sequences of *Phaseolus vulgaris AG* (Phvul.006G169600), *Medicago truncatula AG* (XP\_013462626.1), *Arachis hypogaea AG* (XP\_025640343.1), *Malus domestica AG2* (MDP0000250080), *Malus domestica AG1* (MDP0000324166), *Vitis vinifera AG* (NP\_001268097.1), *Primula vulgaris AG* (AZZ09252.1), *Sesamum indicum AG* (AIS82595.1), *Nicotiana tabacum AG* (NP\_001312829.1), *Populus trichocarpa AG1* (AAC06237.1), *Theobroma cacao AG* (XP\_007025252.1), *Citrus clementina AG* (XP\_006449577.1), *Citrus sinensis AG* (XP\_024950555.1) and *Arabidopsis thaliana AG* (NP\_001328877.1). Values at nodes indicate bootstrap percentages.



**Figure 2.** Relative expression levels of *MdAG1* and *MdAG2* in various tissues of *Malus domestica* 'Golden Delicious'. Expression levels of *MdActin* were used as an internal reference. Data represent means of three biological replicates  $\pm$  SE. Bars with different letters are significant different at *p* < 0.05 based on one-way analysis of variance (ANOVA).

## 2.2. Cloning and Analysis of the Second Introns of MdAG1 and MdAG2

We cloned the sequences of the second introns of *MdAG1* and *MdAG2* and named them as *MdAG11* (MN403053) and *MdAG21* (MN401764), respectively. The two sequences showed 87% identity and were 4,239 bp and 4,236 bp in length, respectively. We further analyzed the putative *cis*-regulatory elements of both intron sequences using PlantCARE [29]. The distributions of some known *cis*-regulatory elements in both intron sequences are shown in Figure 3. Both *MdAG11* and *MdAG21* have core *cis*-regulatory elements, such as TATA-box and CAAT-box elements. For instance, *MdAG11* has 25 TATA boxes and 24 CAAT boxes, while *MdAG21* has 30 TATA boxes and 28 CAAT boxes. In addition, we identified a series of hormone- and environmental-signal-related *cis*-regulatory elements, including three light-responsive elements (GT1 motif, AAAC motif, and G box), one circadian control element

(circadian), one drought-responsive element (MBS), one low-temperature-responsive element (LTR), three hormone-related *cis*-regulatory elements (GARE motif, TGA element, and TGACG motif), and one leafy (LFY)-binding site. The LFY-binding site is reported to bind with *AG* second-intron sequences, a process that plays important roles in maintaining *AG* expression in floral organs [30–32]. We identified one LFY-binding site in *MdAG11* and two in *MdAG21*, and they were also distributed differently within the two intron sequences. These differences may result in different expression patterns when the introns act as promoters.



**Figure 3.** Distributions of ten types of predicted cis-regulatory elements in the second introns of *MdAG1* and *MdAG2*. The different distribution modes may cause different activities when these are used as promoters.

# 2.3. Floral-Specific Expression of rMdAG2I::GUS in Tobacco

To characterize the promoter activity of the two intron sequences, we fused each sequence with the  $\beta$ -glucuronidase (*GUS*) gene. It has been reported that the directions of intron sequences may direct different expression patterns [21]. Therefore, we created four different constructs by fusing the two intron sequences with *GUS* in both forward and reverse directions, creating the constructs *fMdAG11::GUS*, *rMdAG11::GUS*, *fMdAG21::GUS* and *rMdAG21::GUS* (Figure 4).



**Figure 4.** Gene constructs used for tobacco and *Malus* transformation. *35S*, cauliflower mosaic virus 35S gene promoter sequence; *GUS*, coding sequence of the β-glucuronidase gene; *tNos*, nopaline synthase terminator; *fMdAG1I*, forward orientation of the second intron of *Malus AGAMOUS1* (*MdAG1*); *rMdAG1I*, reverse orientation of the second intron of *MdAG2*, *forward* orientation of the second intron of *MdAG2*, *forward* orientation of the second intron of *MdAG2*, *forward* orientation of the second intron of *MdAG2*, *forward*, *fo* 

We first tested these constructs in tobacco. An *Agrobacterium*-mediated stable transformation of tobacco with the four constructs produced 28, 23, 31 and 20 lines for *fMdAG11::GUS*, *rMdAG11::GUS*, *fMdAG21::GUS*, and *rMdAG21::GUS*, respectively. We also produced eight 355::GUS transgenic tobacco

lines. Next, we assayed the leaf, stem and flower tissues of the transgenic tobacco or non-transgenic control by histochemical staining for GUS activity. All the tested tissues exhibited GUS activity in the eight *35S::GUS* lines (Figure 5A–C and Table 1), whereas none of these tissues had GUS activity in the non-transformed control (Figure 5D).



**Figure 5.** GUS staining in tobacco transformed with the four artificial promoters and *355::GUS* control. (**A–C**) GUS expression using *355::GUS* as a positive control. All organs are stained. (**D**) GUS expression in non-transformed flowers. None of the plant was stained. (**E–G**) GUS expression of tobacco tissues transformed with *fMdAG11::GUS* (**E**), *rMdAG11::GUS* (**F**), and *fMdAG21::GUS* (**G**), respectively. Flowers, leaves and stems all show different degrees of blue color. (**H–I**) Two lines of *rMdAG21::GUS* transgenic tobacco. Only flowers are stained.

For the *MdAG1* intron sequence, both the forward and reverse promoters exhibited *GUS* expression patterns similar to those of the *35S::GUS* plants, with GUS activity in all tested tissues. However, *fMdAG11::GUS* caused stronger GUS activity than *rMdAG11::GUS*, indicating that the direction of the intron sequence did affect the activity of the artificial promoter (Figure 5E,F). Of the two promoters derived from the *MdAG2* intron sequence, the forward promoter had GUS activity in both vegetative and floral organs (Figure 5G), whereas the reverse promoter had GUS activity only in floral organs, and not in vegetative organs (Figure 5H,I), suggesting a strong likelihood that *rMdAG21::GUS* governs flower-specific expression of target genes in *Malus*. The difference in expression patterns between the forward and reverse directions of the two intron-sequences demonstrated that their activities are largely dependent on their orientation, which is consistent with our previous findings for the poplar *AG* gene [21]. The direction difference presumably leads to differences in the distances between the *cis*-regulatory sequences and transcription start sites, thus resulting in different expression patterns. In support of this, *MdAG2I* has one more LFY-binding site than *MdAG1I* at the 5' end (Figure 3). Also, we identified only AAAC motif and TGA elements in *MdAG1I*, suggesting that *MdAG1I* may be more easily affected by environment stimuli.

Vectors	No. of Transgenic I	Lines	Plant Tissues						
	0	Stem	Leaf	Sepal	Petal	Stamen	Carpel	I	
35S::GUS	8	+	+	+	+	+	+	No	
fMdAG11::GUS	28	+	+	+	+	+	+		
rMdAG11::GUS	23	+	+	+	+	+	+	- No	
fMdAG2I::GUS	31	+	+	+	+	+	+		
rMdAG21::GUS _	Type I 1	- 11	_	+	+	+	+	Yes	
	Type II	9 –	_	_	_	+	+		

**Table 1.** Characterization of floral specificity of *fMdAG11::GUS*, *rMdAG11::GUS*, *fMdAG21::GUS*, and *rMdAG21::GUS* transgenic tobacco.

+, detectable GUS activity; –, no detectable GUS activity.

Interestingly, *rMdAG21::GUS* lines exhibited two different patterns of *GUS* expression in flower organs (Table 1). Eleven of the 20 lines (Type I) showed GUS activity in all floral organs (sepals, petals, stamens and carpels), while the remaining nine lines showed GUS activity only in stamens and carpels (Type II). Those two patterns were probably due to variable copy numbers or different transgene insertion sites among different lines. Both expression patterns would be useful in practice for breeding purposes. For instance, to produce flowerless plants without any floral organs by fusing a promoter with a toxin gene, we would need a flower-specific promoter with activity in all floral organs as Type I [21]. However, if we planned simply to produce sterile plants with normal petal development for ornamental purposes, the Type II expression pattern would be needed [33–35].

#### 2.4. Optimization of Agrobacterium-Mediated Transient Transformation Conditions for Malus Flowers

We further tested the four constructs in two *Malus* species. Because there is no well-established stable transformation protocol for most *Malus* species, we employed *Agrobacterium*-based transient transformation to test all four constructs. The transient transformation method has the advantages of being simpler and faster than the stable transformation [36–39] and is regularly used in both *Arabidopsis* and tobacco to characterize the subcellular localization of functional genes [40–42]. Additionally, it has been reported that combining the clustered regularly interspersed short palindromic repeat (CRISPR)/Cas9 system and *Agrobacterium*-mediated transient transformation makes it possible to produce a target mutant with no integration of transgenes, which can be very important to accelerate the application of CRISPR technology in asexually-propagated woody plants [43–46]. In *Malus* species, especially *M. domestica*, *Agrobacterium*-mediated transient transformation has been widely used in studying fruit traits because it allows quick testing of the functions of genes associated with fruit traits, avoiding the lengthy periods associated with stable transformation methods because of the long juvenile phase of these species [47]. However, although well established for *Malus* fruit, *Agrobacterium*-mediated transient transformation is not equally successful in different sample types in *Malus* species. Therefore, we first optimized the *Agrobacterium*-mediated transient transformation system for flower tissues.

First, because the efficiency of T-DNA delivery into the host cell is one of the most important factors affecting Agrobacterium-mediated transient transformation, we included a vacuum infiltration step in our transient transformation system. Vacuum infiltration is reported to enhance Agrobacterium infection in some species, such as Arabidopsis, cotton and wheat [48], by improving the penetration efficiency of Agrobacterium cells into target plant tissues. We optimized the vacuum infiltration procedure using a 35S::GUS construct in one representative ornamental Malus species, Malus micromalus (midget crabapple). Different plant tissues may have different tolerances of vacuum pressure due to the major variations in their cell walls and intercellular spaces. Moreover, short vacuum times may not achieve ideal results, while long vacuum times may damage the tissues [49]. Therefore, to achieve an appropriate vacuum time for flower tissues, we tested a gradient of vacuum times: 2.0 min, 3.0 min and 4.0 min, corresponding to vacuum levels of -70 kPa, -80 kPa and -85 kPa, respectively. We also observed that the length of the vacuum application affected the transformation efficiency. Therefore, we tested the results of maintaining the samples at each pressure for four different time periods: 1.0 min, 1.5 min, 2.0 min and 3.0 min. The flowers sustained more serious damage at increased vacuum levels (Table 2); in particular, the -85 kPa vacuum level was unacceptable because it caused relatively serious damage. Maintaining samples under relatively low vacuum pressure (-70 kPa) for a long time improved the staining efficiency. At -80 kPa, a large increase of the treatment time also caused tissue damages. Ultimately, we decided that a -80 kPa vacuum maintained for 1.5 min was ideal, and we used this for later experiments. By employing a modified setting, we were also able to detect GUS activity in stem materials with the 35S::GUS construct, suggesting that, besides the floral organs, the optimized protocol is also available for stem tissues.

Vacuum Time (min)	Soaking Time (min)	Condition of Treated Flowers
2.0	1.0	No damage, No staining
2.0	1.5	No damage, Blue staining
2.0	2.0	No damage, Blue staining
2.0	3.0	No damage, Blue staining
3.0	1.0	No damage, No staining
3.0	1.5	No damage, Blue staining
3.0	2.0	Slightly damaged
3.0	3.0	Damaged
4.0	1.0	Damaged
4.0	1.5	Damaged
4.0	2.0	Damaged
4.0	3.0	Damaged

**Table 2.** Optimization of conditions for *Agrobacterium*-mediated transient transformation of *Malus* 

 flowers using 35S::GUS construct.

#### 2.5. rMdAG2I::GUS Confers Flower-Specific Expression in Malus

Next, we used the optimized transformation conditions for further experiments with the four promoter constructs. We were able to detect GUS activity in all tested organs for 35S::GUS-transformed flowers (Figure 6A), while, none GUS activity was detected for the non-transformed flowers (Figure 6B). In the *fMdAG11::GUS* flowers, we detected GUS activity in both vegetative (stem) and floral organs (sepal, petal, stamen, pistil), which was consistent with the results in tobacco (Figure 6C, Table 3). In *rMdAG11::GUS* flowers, we detected GUS activity in stem and stamen, but not in sepal, petal, or pistil (Figure 6D, Table 3). Notably, in tobacco, *fMdAG11::GUS* produced higher activity than *rMdAG11::GUS* in all instances; therefore, the undetectable activity of *rMdAG11::GUS* in some flower organs in *M*. micromalus may be due to its global low activity. In fMdAG21::GUS flowers, we detected activity in stems and flower organs (Figure 6E, Table 3), which is similar to the expression pattern of *fMdAG11::GUS*. Because those three vectors had GUS activity in stems, they are not suitable for use as flower-specific promoters in *Malus*. Interestingly, we also detected GUS activity in the receptacles of *fMdAG11::GUS*, fMdAG21::GUS and rMdAG21::GUS flowers (Table 3). The reason that the rMdAG11::GUS flowers exhibited no GUS activity in receptacles may be its global low activity. Based on the gene expression data in Figure 2, the expression pattern of both AG genes in fruits may undergo dynamic changes. Our results suggest that the second introns of Malus AG genes may play some role in regulating AG expression during fruit development. For *rMdAG21::GUS*, we did not detect GUS activity in stems, but we did detect GUS activity in all tested floral organs (sepal, petal, stamen and pistil) (Figure 6F, Table 3). Therefore, *rMdAG21::GUS* has strong potential as a flower-specific promoter in *Malus*.

**Table 3.** Characterization of floral specificity of *fMdAG11::GUS*, *rMdAG11::GUS*, *fMdAG21::GUS* and *rMdAG21::GUS* in *Malus micromalus* with *Agrobacterium*-mediated transient transformation.

Organs Vectors	Stem	Sepal	Petal	Stamen	Pistil	Receptacle
35S::GUS	+	+	+	+	+	+
fMdAG11::GUS	+	+	+	+	+	+
rMdAG1I::GUS	+	-	-	+	-	-
fMdAG2I::GUS	+	+	+	+	+	+
rMdAG2I::GUS	-	+	+	+	+	+

+, detectable GUS activity; -, no detectable GUS activity.



Figure 6. GUS staining of *Malus micromalus* with *Agrobacterium*-mediated transient transformation.
(A) 35S::GUS-transformed flowers (positive control). Stem and flower show blue staining.
(B) Non-transformed flowers (negative control) have no blue staining. (C–F) *fMdAG11::GUS-*(C) *rMdAG11::GUS-* (D) *fMdAG21::GUS-* (E) and *rMdAG21::GUS-* (F) transformed flowers show blue staining.

We also tested our optimized transformation protocol with one *M. domestica* cultivar, 'Golden Delicious'; unfortunately, the results were poor, suggesting that *Agrobacterium*-mediated transient transformation is species or material dependent. Thus, we may need to modify this transformation protocol for each new species or cultivar. Given this result, we next turned our attention to an alternative transient transformation method, particle bombardment. Although costly, this method has some important advantages, such as the absence of limitations on species or sample types [50–52]. Therefore, we used particle bombardment to test the flower specificity of *rMdAG21::GUS* in 'Golden Delicious', using *35S::GUS* as control. With *35S::GUS*, we detected GUS activity in both vegetative organs (leaf and stem) and floral organs (sepal, petal, stamen, pistil and receptacle) (Table 4). Meanwhile, with *rMdAG21::GUS*, we did not detect any GUS activity in vegetative organs, but we did detect GUS activity in all floral organs tested (Figure 7 and Table 4). These results further confirm the floral specificity of *rMdAG21::GUS* in *Malus* specie.

**Table 4.** Characterization of the activity of *rMdAG21* in 'Golden Delicious' through particle bombardment.

Organ	Stem	Leaf	Sepal	Petal	Stamen	Pistil	Receptacle
35S::GUS	+	+	+	+	+	+	+
rMdAG2I::GUS	_	_	+	+	+	+	+

<sup>+,</sup> detectable GUS activity; –, no detectable GUS activity.



**Figure 7.** GUS staining of 'Golden Delicious' transformed with *rMdAG21::GUS* through particle bombardment. (**A**) Flower organs with blue staining. RE, receptacle; SE, sepal; PE, petal; ST, stamen; PI, pistil. (**B**) Stem without blue staining.

In summary, we have successfully identified two *AG* paralogs (*MdAG1* and *MdAG2*) from *Malus*. By fusing the second-intron sequences of these two genes to the minimal 35S promoter in both forward and reverse directions, we build four promoter constructs. Next, we demonstrated the flower specificity of the *rMdAG21* promoter through the stable transformation of tobacco plants. We further verified its flower specificity in the *Malus* species by Agrobacterium-mediated transient transformation and particle bombardment. This newly designed *rMdAG21* promoter thus has great potential as a means for the genetic modification of flower traits in the *Malus* species.

## 3. Materials and Methods

## 3.1. Phylogenetic Analysis

Phylogenetic analysis was conducted using MEGA 6.0 software [53]. The tree was compiled using the neighbor-joining method. Statistical reliabilities of the internal branches were assessed using 1000 bootstrap replicates. Values at nodes indicate bootstrap percentages [54].

## 3.2. RNA Extraction and qRT-PCR

The vegetative organs (roots, stems and leaves) and reproductive organs (receptacles, sepals, petals, stamens, pistils and fruits) of 'Golden Delicious' apples were prepared for RNA extraction. Total RNA of each sample was extracted using the cetyl trimethyl-ammonium bromide (CTAB) method. RNA quality and concentration were determined using a NANODROP 2000 spectrophotometer (Thermo Scientific, Waltham, MA, USA). The cDNA was synthesized using MLV reverse transcriptase (TakaRa Bio Inc., Dalian, China). We performed quantitative real-time PCR (qRT-PCR) analysis using the SYBR premix Ex Taq Kit (TakaRa Bio Inc., Dalian, China) in a 7500 real-time PCR system (Applied Biosystems Foster City, CA, USA).

The primer pair AG1-F (5'-GAGAGGTACAAGAAGGCAAGTGC-3') with AG1-R (5'-CTCTTC AGGTCCTTGACAGACATAC-3') was used for the analysis of *MdAG1* transcripts, and the pair AG2-F (5'-ATGCCTATCAAGGACCTGAAGAG-3') with AG2-R (5'-TAGTTC CGAGAGTCGTATGGCT-3') was used for analysis of *MdAG2* transcripts. The primer pair Actin-F (5'-TGACCGAATGAGCAA GGAAATTACT-3') with Actin-R (5'-TACTCAGCTTTGGCAAT CCACATC-3') was used to amplify the internal reference gene, *MdActin* [47]. Data were analyzed using the 2<sup>- $\Delta\Delta$ CT</sup> method [55]. The gene expression in each sample was calculated using the expression level of *MdActin* in the same sample. Each qRT-PCR reaction was performed with three biological replicates, and each sample was analyzed in triplicate (technical replicates).

## 3.3. Cloning of MdAG1 and MdAG2 Second-Intron Sequences

Genomic DNA was extracted from the leaves of 'Golden Delicious' using a modified CTAB method [56]. Approximately 200 ng genomic DNA was used as templates for PCR amplification of the second-intron fragments from two *MdAG* genes. PCR amplification of the *MdAG1* second intron was performed using the primer pair AG1I-F (5'-CGCCTCTATGAGTATGCCAAC-3') and AG1I-R (5'-CTTCTTGTACCTCTCAATTGTTCCT-3') under the following conditions: 5 min at 95 °C, followed by 35 cycles of 30 s at 94 °C, 30 s at 58 °C, and 2 min at 72 °C, with a final extension of 7 min at 72 °C. PCR amplification of the *MdAG2* second intron was performed with the primer pair AG2I-F (5'-GAGTATGCCAACAACAAGGTAAT-3') and AG2I-R (5'-TGCTTTAACACTGCAACAATATGG-3') under the following conditions: 5 min at 95 °C, followed by 35 cycles of 30 s at 94 °C, 30 s at 58 °C, and 2 min at 72 °C. The amplified fragments were inserted into the pEASY-Blunt Cloning Kit, and 10–12 clones were sequenced for verification of their sequence identity.

#### 3.4. Plasmid Construction

Four chimeric promoters were created by fusing the intron sequence of either *MdAG1* or *MdAG2* to a 60-bp minimal 35S promoter in either forward or reverse direction. Those four artificial promoters were inserted upstream of the *GUS* coding sequence to create four plasmids: *fMdAG1::GUS*, *rMdAG1::GUS*, *fMdAG2::GUS*, and *rMdAG2::GUS*. A control vector was also created containing the *GUS* coding sequence driven by a constitutively active 35S promoter (*35S::GUS*).

## 3.5. Stable Transformation of Tobacco

Each of the four plasmids was introduced into *Agrobacterium tumefaciens* EHA105, which was subsequently used for transformation of tobacco (*Nicotiana benthamiana*) [57]. Leaf discs of tobacco ( $0.5 \text{ cm} \times 0.5 \text{ cm}$ ) were incubated for 20 min with *A. tumefaciens* ( $OD_{600} = 0.6$ ) and then transferred to the Murashige and Skoog (MS) medium without hormone and antibiotics for two days of co-cultivation. After that, leaf explants were moved to a regeneration medium containing 3 mg/L 6-benzylaminopurine (6-BA), 0.2 mg/L 1-naphthaleneacetic acid (NAA), 300 mg/L cefotaxime, and 50 mg/L kanamycin. Putative transgenic shoots were moved to rooting medium containing 300 mg/L cefotaxime and 20 mg/L kanamycin. The rooting shoots were transferred to soil and grown in a greenhouse.

#### 3.6. Vacuum Infiltration of Malus Micromalus Flowers

Agrobacterium tumefaciens EHA105 harboring each of four created constructs was prepared. Flowers of *M. micromalus* were immersed in the *Agrobacterium* suspension ( $OD_{600} = 1.0$ ) and placed in a vacuum chamber. A total of 12 treatments with three vacuum treatment times (2 min, 3 min, 4 min) and four soaking times (1 min, 1.5 min, 2 min, 3 min) were used for orthogonal tests. After treatments, the samples were cultured on MS medium for two days and then subjected to GUS staining.

# 3.7. Particle Bombardment of 'Golden Delicious' Apple Flowers

To prepare the gold suspension, a mixture of gold powder and ethanol (60 mg:1 mL) was shaken for 1 min and centrifuged at 10,000 rpm for 10 s. The pellets were then resuspended in sterilized water. 50  $\mu$ L of gold suspension, 20  $\mu$ L 0.1 M spermidine, 50  $\mu$ L 2.5 M CaCl<sub>2</sub>, and 2.5  $\mu$ g DNA were used for preparing the gold/DNA mixture. After having been shaken for 3 min, the mixture was centrifuged at 10,000 rpm for 20 s. The pellets were rinsed by ethanol for two times and then re-suspended by 60  $\mu$ L ethanol, and 10  $\mu$ L was used for each particle bombardment. The bombardment was repeated six times for each sample. The treated samples were cultured on MS medium for two days before GUS staining.

## 3.8. GUS Staining Assays

GUS assays were performed according to a previously described protocol with modifications [21]. Samples were incubated in X-Gluc solution (100mM potassium phosphate buffer, 10 mM Na<sub>2</sub>EDTA, 0.5 mM K<sub>4</sub>Fe (CN)<sub>6</sub>, 0.1% Triton X-100, and 1 g/L X-Gluc, pH 7.0) at 37 °C. After 24 h of incubation, the samples were decolorized with alcohol three to five times before being photographed.

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