CRISPR/Cas9-mediated editing of 1-aminocyclopropane-1-carboxylate oxidase1 enhances Petunia flower longevity

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Introduction

Petunias, widely used as a bedding plant in the floricultural industry, have become increasingly popular due to their diversity of different flower shapes and colours. Generally, petunias exhibit excellent flower longevity because they continuously produce new flowers over an extended period. However, newly produced individual flowers exhibit rapid senescence in the mother plants. Because petunias are ethylene-sensitive, their flower senescence is associated with an increase in ethylene production (Woltering and Van Doorn, 1988; Tang et al., 1993; Tang and Woodson, 1996; Huang et al., 2007). Hence, it is necessary to find an effective way to reduce ethylene production.

Ethylene is derived from methionine, which is converted to S-adenosylmethionine (SAM or AdoMet) by SAM synthetase and then to 1-aminocyclopropane-1-carboxylic acid (ACC) by ACC synthetase, before ACC oxidase (ACO) finally converts ACC to ethylene (Yang and Hoffman, 1984). In petunias, it has been reported that PhACO1, PhACO3 and PhACO4 encode ACO and mutant lines, regardless of mutant type (homozygous or monoallelic), exhibited significantly reduced ethylene production and enhanced flower longevity compared with wild-type. Flower longevity and the reduction in ethylene production were observed to be stronger in homozygous plants than in their monoallelic counterparts. Additionally, the transmission of the edited gene to the T2 (lines 6 and 36) generation was also confirmed, with the results for flower longevity and ethylene production proving to be identical to those of the T2 mutant lines. Overall, this study increases the understanding of the role of PhACO1 in petunia flower longevity and also points to the CRISPR/Cas9 system being a powerful tool in the improvement of floricultural quality.

Summary

The genes that encode the ethylene biosynthesis enzyme 1-aminocyclopropane-1-carboxylate oxidase (ACO) are thought to be involved in flower senescence. Hence, we investigated whether the transcript levels of PhACO genes (PhACO1, PhACO3 and PhACO4) in Petunia cv. Mirage Rose are associated with ethylene production at different flowering stages. High transcript levels were detected in the late flowering stage and linked to high ethylene levels. PhACO1 was subsequently edited using the CRISPR/Cas9 system, and its role in ethylene production was investigated. PhACO1-edited T0 mutant lines, regardless of mutant type (homozygous or monoallelic), exhibited significantly reduced ethylene production and enhanced flower longevity compared with wild-type. Flower longevity and the reduction in ethylene production were observed to be stronger in homozygous plants than in their monoallelic counterparts.

The deletion or editing of target genes can be achieved using three genome editing methods: zinc-finger nucleases (ZFNs), transcription activator-like effector nucleases (TALENs) and clustered regulatory interspaced short palindromic repeat (CRISPR/Cas9) sequences. Of these, the CRISPR/Cas9 system has received significant recent attention because of its high specificity for the editing of target genes, its low cost and the simplicity of its design (Cong et al., 2013; Feng et al., 2013; Shan et al., 2013). This system induces site-specific double-strand breaks (DSBs) at a target site upstream of the protospacer-adjacent motif sequence (PAM) within the genome, followed by a DSB repair mechanism consisting of homologous recombination (HR) or non-homologous end-joining (NHEJ). HR leads to the accurate reconstruction of the original sequences using an undamaged homologous sequence or externally supplied donor DNA template to repair the DSBs, while NHEJ repairs the DSBs regardless of homology, leading to insertions or deletions (indels; Hsu et al., 2014). In eukaryotic cells, DSBs are preferentially repaired using NHEJ, thus providing a promising editing strategy for research on plant.
functional genomics and crop improvement (Lieber, 2010; Pan et al., 2016). Recently, this system has been successfully utilized to generate mutagenesis in various plant species such as tomatoes, grapes and rice (Brooks et al., 2014; Miao et al., 2013; Pan et al., 2016; Wang et al., 2017b), and the mutant allele scan also be stably transmitted to the subsequent generation. Hence, the use of CRISPR/Cas9 to modify the genetic loci associated with ethylene biosynthesis in petunias would hold great promise, even though this technique has not been widely employed in ornamental plants.

In this study, we characterized the expression patterns of the ethylene biosynthesis genes PhACO1, PhACO3 and PhACO4 in petunia petals at different stages of development. The PhACO1 gene, which is highly expressed during the flowering period, was edited with the CRISPR/Cas9 system in order to generate PhACO1-edited mutants that exhibited lower ethylene production and subsequently greater flower longevity.

Results

Expression patterns of ethylene biosynthesis genes at different flowering stages

The differential expression of the ethylene biosynthesis genes (PhACO1, PhACO3 and PhACO4) was observed at different flowering stages of Petunia cv. Mirage Rose. The expression patterns were similar for PhACO1 and PhACO3; their transcript levels were relatively low during the flower budding stages (stages 1 and 2), started to slightly increase during the early blooming stages (stages 3 and 4) and then continuously increased during the fully blooming stages (stages 5–7) (Figure 1a). A similar trend was observed for PhACO4, but, compared to PhACO1 and PhACO3, higher transcript levels were first observed during the early blooming stage (stage 4) and started to fall during stage 7 (Figure 1d). Generally, the transcript levels of PhACO1 were higher than those of PhACO3 and PhACO4 for all flowering stages (stages 1–7), whereas those of PhACO4 were the lowest. The ethylene levels detected during the different flowering stages were likely to be associated with the expression patterns of PhACO1, PhACO3 and PhACO4 because ethylene levels were relatively higher during the blooming stages (stages 5–7) than in the early blooming stages (stages 3 and 4; Figure 1e), with the lowest levels observed during the budding stages (stages 1 and 2).

In vivo validation of the designed Cas9-sgRNAs for PhACO1 in petunia protoplasts

To generate PhACO1-edited petunia plants, we designed two specific sgRNAs using Cas-Designer equipped in RGEN tools (http://rgenome.net) and CRISPR-P web tool (http://cbi.hzau.edu.cn/cgi-bin/CRI SPD). To assess the genome editing efficiency of designed sgRNAs (sgRNA1 and sgRNA2) targeting PhACO1, petunia protoplasts were transfected with preassembled Cas9–sgRNA ribonuclease protein (RNP) complexes via polyethylene glycol (PEG)-mediated delivery. After delivering the Cas9 RNP complexes, we incubated the transfected protoplasts at room temperature for 24 h to induce the DSBs at the PhACO1 loci. Fragments surrounding the targeted sequences of the PhACO1 were amplified using PCR with the specific primers (Table S2) and analysed with targeted deep sequencing to detect the insertions/deletions (indels) at the expected positions, three base pairs (bp) upstream of an NGG PAM. Mutations were readily detected at all target sites (Figure 2b) after 24 h of incubation, with the indel frequency for sgRNA1 observed to be higher than for sgRNA2 (up to 4.85% and 2.21%, respectively).

Analysis of the genotype in PhACO1-edited petunias

PhACO1, which had the strongest influence of the three target genes on ethylene production in Petunia cv. Mirage Rose, was targeted by Cas9:sgRNA1, which exhibited a high indel percentage in protoplast transient assays. The pBATC:sgRNA1 construct was delivered into petunia leaf discs using Agrobacterium-mediated transformation, while discs transformed with an empty vector (pBAIC) were used as the WT. The construct-transformed leaf discs started to form calluses after a few weeks of culture, and the calluses then produced shoots in phosphinothricin (PPT)-containing medium. The multiple shoots originating from the callus were regarded as independent transgenic lines. Delivery of the construct in the plants was screened using PCR analysis (Figure S1).

To evaluate the types of mutation at the PhACO1 locus of the transgenic plants, PCR amplification and targeted deep sequencing were conducted. Mutant lines showing different indel patterns were observed when pBATC:sgRNA1 was delivered but were not observed in the WT. According to the results shown in Table 1, a mutation frequency of 31.5% was observed following the delivery of pBATC:sgRNA1, with homozygous, monoallelic mutants and chimeric mutants accounting for 2.5%, 15.0% and 82.5% of the mutants, respectively. Homozygous mutants were characterized by a 10-bp deletion 4 bp upstream of the PAM sites (Figure 3a, line 91[1]), while other mutant lines exhibited different indel patterns at different cleavage sites (Figure 3a). Of the chimeric mutants, we chose lines 91(2) and 109 as representative lines for the determination of flower longevity and ethylene production based on their high indel read percentage. The indel patterns of the other chimeric mutants are presented in Figure S2.

Analysis of the phenotype in PhACO1-edited petunias

To determine flower longevity, flowers of the mutant lines, including the WT line, that opened on the same day, were labelled in the mother plants, and they were allowed to bloom until one-third of the petals exhibited in-rolling. Of the investigated lines, the flower longevity of line 109 was found to be the highest, followed by lines 91(1), 129(2) and 121(1), while the other lines also exhibited longer flower life than the WT (Figures 3b and 4), which were in accordance with the results for the ethylene levels in the mutant lines. The production of ethylene continuously increased until the end of flowering, though the ethylene levels detected in the WT were notably higher than those of the mutant lines in both the petals and pistils 5 days after full blooming (Figure 5a,b). The lower production of ethylene in lines 109, 121(1), 129(2) and 91(1) was associated with lower transcript levels of the target PhACO1 gene, which had the strongest influence of the three target genes on ethylene production in Petunia cv. Mirage Rose, was analysed using Sanger sequencing. Compared with the sequence (23 bp) of PhACO1-sgRNA1, there were six mismatched regions in PhACO3.
and seven mismatched regions in PhACO4 (Figure 7a). However, Sanger sequencing revealed that there were no mutations in this region of the homologous genes PhACO3 and PhACO4 in any of the PhACO1-edited petunias (Figure 7b), thus proving that there was no off-target effect.

Inheritance of PhACO1 editing in T1 petunias

T1 seedlings of all mutant lines (except line 91-1) were obtained from self-pollination in the greenhouse. T1 seedlings from the lines were randomly selected and sequenced to analyse sexual transmission, with different patterns of segregation observed in the lines (Table 2). In addition, their indel patterns were also described (Figure 8). For further analysis of flower longevity and ethylene production, T1 line 36, consisting of 6% homozygous (1/15), 66% monoallelic (10/15) and 26% WT (4/15), and line 6, consisting of 20% homozygous (3/15), 60% monoallelic (9/15) and 20% WT (3/15), were selected. When flower longevity and ethylene production were evaluated for these lines, the results differed significantly depending on the mutant type. Longer flower longevity was observed in the homozygous mutants, followed by the monoallelic and WT (i.e. homozygous > monoallelic > WT), an outcome that was linked to a reduction in ethylene levels (Figure 9a–f). However, the characteristics of the vegetative and floral organs did not significantly differ between the mutant types (WT, homozygous and monoallelic) of lines 6 and 36 (Table S6). Taken together, it is obvious that the editing of...
the \textit{PhACO1} gene in \textit{Petunia} cv. Mirage Rose significantly reduced ethylene production and improved flower longevity, and stable results were also observed in \textit{ACO1}-edited T1 seedlings without affecting the morphology of the vegetative and floral organs.

\textbf{Discussion}

The editing of target genes using CRISPR/Cas9 systems has become popular in recent years due to their ability to induce target mutagenesis with more precision and higher efficiency compared to other genome editing technologies. Over the past 5 years, successful genome editing using this technology in plant species such as maize, wheat, tomatoes and rice has been reported (Li \textit{et al.}, 2018; Svitashev \textit{et al.}, 2016; Zong \textit{et al.}, 2017). Petunia cv. Mirage Rose (\textit{P. hybrida}) is a bedding plant that is widely used in the landscape industry. As these petunias are typically planted in open landscape fields, the reduction in flower longevity caused by ethylene production cannot be
controlled using chemicals that block ethylene biosynthesis. In this study, therefore, we attempted to edit the primary ethylene biosynthesis gene involved in the production of ethylene in Petunia cv. Mirage Rose using the CRISPR/Cas9 system.

Initially, we characterized the expression patterns of the ethylene biosynthesis genes PhACO1, PhACO3 and PhACO4 during different flowering stages (stages 1–7). The transcript levels of the genes were significantly higher during the fully open stages (stages 5–7) than during the initial flowering stages, which are in line with results reported by Tang et al. (1994), who observed an increase in ACO mRNA in petunia petals during flower senescence. In the present study, the transcript levels of PhACO1 in the petals continuously increased and were significantly higher than those of PhACO3 and PhACO4, indicating that the transcript levels of PhACO1 are more strongly linked to the production of ethylene during the flowering stages. Huang et al. (2007) reported the overexpression of the antisense BoACO1 gene (from broccoli) in petunias, whose sequence is 90% homologous to PhACO1, observing the reduction in ethylene production and delayed flower senescence. Therefore, we concluded that PhACO1 is more closely involved in the production of ethylene in this cultivar than the other two genes.

To edit PhACO1, we designed two sgRNAs (sgRNA 1 and sgRNA2) to target PhACO1, and transfected petunia protoplasts with a pBAtC : sgRNAs ribonuclease protein (RNP) complex. We found that sgRNA1 produced higher indel rates compared with sgRNA2; thus, the former was used to edit PhACO1. As expected, a high mutation frequency (31.5%) with different indel patterns and mutant types (i.e. homozygous, monoallelic and chimera) was observed following the delivery of pBAtC : sgRNA1 using Agrobacterium-mediated transformation, and the PhACO1-edited mutant lines improved flower longevity compared with WT plants by reducing ethylene levels, regardless of the mutant type. Of the mutant lines, the homozygous line (line 91-1) and the chimeric line (line 109) significantly reduced the production of ethylene in both petals and pistils, which was linked to the down-regulation of the target PhACO1 gene. Moreover, the PhACO1-edited mutants did not have an off-target effect on two genes (PhACO3 and PhACO4) with a high level of homology, indicating high precision editing of the target gene.

Of the monoallelic mutant lines, lines 121(1) and 129(2) reduced ethylene levels in petals to a greater extent than did the other monoallelic lines (i.e. lines 6, 36, 133[2] and 178[1]). This is linked to the differences in transcript levels for the target PhACO1 among the mutant lines, with lower transcript levels for PhACO1 in the former group lines than in the latter. It is possible that differences in indel patterns significantly affected the transcript levels of the target gene. From these results, it can be concluded that PhACO1 is strongly involved in the production of ethylene, which influences flower longevity in Petunia cv. Mirage Rose, which is consistent with previous research (Huang et al., 2007).
T1 petunias inherited most of the mutation types from the T0 generation. Similar inheritance patterns have been reported in tomatoes by Pan et al. (2016). The results of the present study support the feasibility of using CRISPR/Cas9 to edit a target gene and to transmit the edited gene to the next generation. This contrasts with the lack of homozygotes and biallelic specimens in the T1 generation of CRISPR/Cas9 transgenic Arabidopsis plants (Feng et al., 2014). It is possible that some of the mutations found in somatic cells were lost because the cells did not contribute to germ-line development (Xu et al., 2015). Of the T1 mutant lines, ethylene production in the petals and pistils of lines 6 and 36 was analysed. The lowest ethylene production was observed in homozygous mutants, followed by monoallelic and WT for both lines, as observed in the T0 plants. In addition, the reduction in ethylene levels was positively associated with an increase in flower longevity for both T1 lines (homozygous > monoallelic > WT). These findings also provide further evidence of the stable involvement of ACO1 in ethylene production in this cultivar. It was also found that editing the PhACO1 gene in petunias using CRISPR/Cas9 did not affect the morphology of the vegetative and floral organs.

To date, the editing of ethylene biosynthesis genes has not been reported for any plant species; thus, to our knowledge, this is the first report to generate a novel cultivar with extended flower longevity using CRISPR/Cas9 technology. Anthocyanin 1 (ANT1; Cermak et al., 2015), phytoene desaturase (SlPDS) and phytochrome-interacting factor (SlPIF4; Pan et al., 2016), which are involved in flavonoid and carotenoid biosynthesis in tomato plants, have been edited using CRISPR/Cas9. In addition, the application of CRISPR/Cas9 in editing the PDS gene in petunias has been reported (Zhang et al. 2016); however, the editing of the PDS gene did not lead to actual improvement in petunia traits because PDS-edited plants are not commercially attractive for the floricultural industry. The present study illustrates that the targeted editing of PhACO1 can be achieved using CRISPR/Cas9 and that PhACO1-edited petunias exhibit significantly extended flower longevity due to a significant reduction in ethylene production. Hence, we believe that CRISPR/Cas9-based ethylene tuning is likely to be an effective strategy for enhancing flower longevity in other floricultural crops.

Conclusion

We investigated the potential roles of PhACO genes (PhACO1, PhACO3 and PhACO4) in the ethylene production of petunia flowers using transcriptional analysis at different stages of flowering. Of the PhACO genes, we revealed that PhACO1 was
most likely to be involved in ethylene biosynthesis. The application of a CRISPR/Cas9 tool in petunias efficiently generated several T0 transformants with different indel patterns. The edited petunias exhibited significantly extended flower longevity by reducing ethylene levels compared with the WT. Furthermore, T1 petunias inherited the editing patterns, reduced ethylene levels and the phenotype of extended flower longevity that were similar to the parental genotype and phenotype. Our study contributes to a better understanding of the underlying role of PhACO1 in ethylene biosynthesis and flower longevity. In addition, it provides evidence for the feasibility of the use of CRISPR/Cas9 to precisely edit PhACO1 and to determine its function; thus, our study can pave the way for the editing of ethylene biosynthesis genes in other ornamental plants, thereby advancing plant biology and the floricultural industry.

Experimental procedures

Plant materials

Petunia (Petunia hybrida cv. Mirage Rose) seeds obtained from the Hanmi seedling company (Korea America Plug Co., Ltd, Gunpo-si, Gyeonggi-do, South Korea) were sown in plug trays filled with a soil-less mixture (Berger Co., Quebec, QC, Canada) in a greenhouse with a day/night temperature of 22°C/18°C, a photoperiod of 16 h and a relative humidity of 70%. After 2 weeks, the germinated seedlings were transferred to individual pots filled with the same soil and grown under the same greenhouse conditions until flowering.

Characterization of the expression patterns of ethylene biosynthesis genes

To investigate the expression patterns of ethylene biosynthesis genes (PhACO1, PhACO3 and PhACO4) in the flowers of Petunia cv. Mirage Rose, petals (approximately 500 mg) from different flowering stages (Figure 1a) were placed in 2-mL tubes immersed in liquid nitrogen. They were then immediately stored at −80°C for RNA extraction. Total RNA extraction and reverse transcription were performed as described by Naing et al. (2017a). The transcript levels of PhACO1, PhACO3 and PhACO4 were measured relative to those of alpha-tubulin gene (reference gene) using the StepOnePlus™ Real-Time PCR system (Thermo Fisher Scientific, Waltham, MA). Relative gene expression was calculated using the quantitative-comparative CT (ΔΔCT) method. The primers and polymerase chain reaction (PCR) conditions for the detected genes are listed in Table S1. The analysis was repeated three times for each stage.

Table 2 Segregation patterns of CRISPR/Cas9-mediated mutagenesis in T1 generation

| Target gene | Line | Zygosity | Genotype | Mutation segregation | Cas9 |
|-------------|------|----------|----------|----------------------|------|
| ACO1        | T0-6 | Monoallelic | i1WT | 3i1i1, 9i1WT, 3WT | All+ |
|             | T0-36 | Monoallelic | d5WT | 1d5d5, 10d5WT, 4WT | All+ |
|             | T0-91 | Chimeric | d4WT | 6d4WT | 5+;1− |
|             | T0-10 | Chimeric | i1d5 | 1i1, 2d5d5, 4i1d5, 1i1WT | All+ |
|             | T0-12 | Monoallelic | d10WT | 7d10d10, 6d10WT | All+ |
|             | T0-133 | Monoallelic | i1WT | 1i1, 6i1WT, 6WT | All+ |
|             | T0-176 | Monoallelic | d2WT | 5d2d2, 6d2WT, 4WT | 10+;5− |

Table 2 continued

| Target gene | Line | Zygosity | Genotype | Mutation segregation | Cas9 |
|-------------|------|----------|----------|----------------------|------|
| ACO1        | T0-109 | Chimeric | d4WT | 6d4WT | 10+;5− |
|             | T0-121 | Chimeric | i1d5 | 1i1, 2d5d5, 4i1d5, 1i1WT | All+ |
|             | T0-133 | Monoallelic | d2WT | 5d2d2, 6d2WT, 4WT | 10+;5− |

+, Cas9 was detected; −, Cas9 was not detected; #, # of nucleotide(s) deleted at the target sites; #: # number of nucleotide(s) inserted at target sites; WT, wild-type sequence without mutations detected at target sites.

Figure 8 Transmission of the PhACO1-edited gene to T1 mutant lines (6 and 36) identified by sequencing. The minus (−) and plus (+) signs indicate the number of nucleotides deleted and inserted at the target sites. WT, wild-type sequence without mutations at the target sites.

Edition of ACO1 gene in Petunia 293

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guide sequences complementary to the different DNA strands of PhACO1 in the exon regions (exon 1: 19–38 and exon 2: 37–59; Figure 2a).

sgRNAs were synthesized via in vitro transcription using T7 RNA polymerase (New England Biolabs, Ipswich, MA) in a reaction buffer (40 mM Tris-HCl, 6 mM MgCl2, 10 mM DTT, 10 mM NaCl, 2 mM spermidine, NTPs and RNase inhibitor, pH 7.9) according to the manufacturer’s protocol (Kim et al., 2014). Transcribed sgRNAs were incubated with DNase I at 37 °C for 30 min to remove template DNA and then purified using PCR purification kits (GeneAll, Seoul Korea).

Protoplast isolation and transient assays

Protoplasts were isolated from the leaves of 5-week-old petunia plants using the valosin-containing protein (VCP) enzyme (Jie et al., 2011). Briefly, the leaves were chopped and digested in 20 mL of enzyme solution at room temperature under gentle shaking (40 rpm) in the dark for 3 h. Following this, the mixture was diluted with an equal volume of W5 solution (154.0 mM NaCl, 125.0 mM CaCl2, 5.0 mM KCl, 5.0 mM glucose, 1.5 mM MES) and filtered through a 100-μm nylon mesh (Falcon, Capitul Scientific, Inc, TX, Austin) to remove undigested leaf tissue (Yoo et al., 2007). Protoplasts were then purified and collected using sucrose gradient centrifugation at 70 g for 7 min before being transferred to W5 solution (Jie et al., 2011).

Recombinant Cas9 protein (30 μg) and in vitro-transcribed sgRNA (80 μg) were mixed and incubated at room temperature for 10 min to generate a ribonuclease protein (RNP) mixture (Woo et al., 2015). Protoplasts (1 × 10^5) in MMG solution (4 mM MES, 0.4 M mannitol and 15 mM MgCl2) were then gently mixed with the RNP mixture, and the RNP mixture was transfected into the protoplasts via PEG-mediated transfection (Woo et al., 2015; Yoo et al., 2007). Briefly, protoplasts (1 × 10^5) were blended with the RNP mixture with an equal volume of PEG solution (40% w/v PEG 4000, 0.2 M mannitol and 0.1 M CaCl2) for 20 min at room temperature, and then, the protoplasts were washed with W5 solution. The transfected protoplasts were incubated in W5 solution at room temperature for 24 h. After 24 h, the protoplasts were collected by centrifugation and the genomic DNA extracted for analysis using targeted deep sequencing.
DNA extraction and targeted deep sequencing

Genomic DNA was extracted from transfected protoplasts using the DNeasy Plant Mini Kit (Qiagen, Hilden, Germany). The target region was amplified using nested PCR primer pairs containing adapter sequences. Following this, the amplicons were labelled with an index sequence (Illumina, Seoul, South Korea) using index PCR primer pairs. The targeted deep sequencing of the index PCR amplicons was conducted using an Illumina MiniSeq. The targeted deep sequencing data were analysed using a Cas-analyzer (Park et al., 2016). The site-specific primer pairs used in this study are listed in Table S2.

Vector construction

We used a pBAtC vector (accession number KU213971) and Aar I-mediated sgRNA cloning system (Kim et al., 2014) for Agrobacterium-mediated transformation in Petunia cv. Mirage Rose. The target sgRNA sequence for ACO1-sg1 was inserted as an annealed oligonucleotide into the AarI-digested pBAtC vector between the AtU6 promoter and sgRNA scaffold and then circularized by incubation with T4 ligase (New England Biolabs, Ipswich) at room temperature for 3 h. An annealed oligonucleotide into the AarI-digested pBAtC vector was amplified using a mixture containing 100 pmol/L of target oligonucleotides under the following conditions: 95 °C for 5 min, a linear gradient from 95 to 25 °C for 70 min and holding at 10 °C. The ACO1 (exon 1) targeting sgRNA (gx20) sequence was transcribed under the control of the Arabidopsis thaliana-U6 promoter, and human codon-optimized Streptococcus pyogenes Cas9 (SpCas9) expression was controlled by the 3SS-promoter.

Agrobacterium-mediated transformation

The transformation of petunia leaf explants was conducted as described by Ai et al. (2017). Briefly, approximately 700 leaf explants were excised from 5-week-old petunia plants grown in the greenhouse and initially pre-cultured in are generation medium. The pre-cultured explants were then infected with Agrobacterium tumefaciens that harboured the plasmid vector. The explants were then dried on sterile filter paper and cultured in the regeneration medium in the dark for 2 days. Following this, the explants were transferred to a selection medium containing 0.5 mg/L PPT. Shoots that showed resistance to PPT were transferred to hormone-free MS medium containing a higher concentration of PPT (1.0 mg/L). Plantlets that produced roots in the PPT-containing (1.0 mg/L) media were completely removed, washed thoroughly with sterile distilled water, transferred to plastic pots containing peat-based soil and grown in a greenhouse under controlled conditions.

Determination of the transcript levels of the PhACO1 gene in the mutants

To determine the transcript levels of the target PhACO1 gene expressed in the PhACO1-edited mutant lines, total RNA extraction and reverse transcription were performed as described above. In addition, the transcript levels of the genes were also measured as described above. The primer and PCR conditions used for the transcriptional analyses are described in Table S4.

Determination of flower longevity and ethylene levels

Flowers of the mutant lines, including the WT line, that opened on the same day, were labelled in the mother plants. The labelled flowers were allowed to bloom until one-third of the petals exhibited in-rolling, and the longevity of each flower was determined. Longevity was assessed using 10 flowers per mutant line with three replications.

To measure ethylene production, petals and pistils (approximately 500 mg or 50 mg) from each mutant line (including WT) were weighed and sampled 1, 3, 5 and 7 days after flowering. They were placed in a 50-mL glass tube, which was then sealed with a rubber septum, and ethylene was detected as described by Naing et al. (2017a, 2017b, 2017c). Three replicates were used for each line.

Sanger sequencing

Total genomic DNA from each T0 mutant plant was used as a template to amplify the target regions of the PhACO1 genes by PCR using the corresponding primer pairs (Table S5). Amplicons were directly sequenced using the Sanger sequencing method (Macrogen, Seoul, South Korea).

Generation of T1 lines and the determination of flower longevity and ethylene levels

The flowers from the T0 mutant lines were self-pollinated in the greenhouse. The seeds obtained were then germinated in vitro in MS basal medium, and the regenerated plants were analysed using PCR, followed by deep sequencing as described above. Of the T1 mutant lines, only lines 6 and 36 were selected to assess flower longevity and ethylene levels using the process described above.

Statistical analysis

The data were analysed using SPSS v. 11.09 (IBM Corporation, Armonk, NY) and presented as the mean of three replicates. Duncan’s multiple range test (DMRT) was used to assess the differences between the means. The significance level was set at $P < 0.05$.

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Conflicts of interest

There are no conflicts of interest among the authors.
Author contributions
C. K. K. and A. H. N. designed the experiments. J. -S. Kim, B.-C. K. and H. K. generated all the constructs. J. X. analysed transcription levels, flower longevity and ethylene production. B.-C. K. performed RNP transient assays for the protoplasts. J. X. generated the transgenic lines. B.-C. K. and S.-J. B. analysed the Sanger sequencing and targeted deep sequencing. B.-C. K. helped in writing some parts of the methods section. A.H.N and J. X. statistically analysed all data. A. H. N. wrote the manuscript. A. H. N. and H. K. revised the manuscript. C. K. K supervised the project.

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Supporting information
Additional supporting information may be found online in the Supporting Information section at the end of the article.
Table S1 Primer sequences used for gene expression analysis using quantitative real-time PCR.
Table S2 Primer sequences used for nested PCR and 2nd round PCR analysis.
Table S3 Primers and PCR conditions used for the detection of the presence of the genes using PCR.
Table S4 Primers used for the transcriptional analysis of PhACO1 in the mutants.
Table S5 Primers used for Sanger sequencing.
Table S6 Comparison of the morphological characteristics of the wild-type and T₁ mutant lines.

Figure S1 Detection of the presence of Cas9-1, Cas9-2, and the Basta gene in the transgenic lines in comparison with plasmid and wild-type specimens using simple PCR.

Figure S2 Scheme for the insertion/deletion patterns of different T₀ mutant lines identified by sequencing. Minus (−) and plus (+) signs indicate the number of nucleotides deleted and inserted at the target sites.