PhCESA3 silencing inhibits elongation and stimulates radial expansion in petunia

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Cellulose synthase catalytic subunits (CESAs) play important roles in plant growth, development and disease resistance. Previous studies have shown an essential role of *Arabidopsis thaliana* CESA3 in plant growth. However, little is known about the role of CESA3 in species other than *A. thaliana*. To gain a better understanding of CESA3, the petunia (*Petunia hybrida*) PhCESA3 gene was isolated, and the role of PhCESA3 in plant growth was analyzed in a wide range of plants. PhCESA3 mRNA was present at varying levels in tissues examined. VIGS-mediated PhCESA3 silencing resulted in dwarfing of plant height, which was consistent with the phenotype of the *A. thaliana* rsw1 mutant (a temperature-sensitive allele of AtCESA1), the *A. thaliana* cev1 mutant (the AtCESA3 mild mutant), and the antisense AtCESA3 line. However, PhCESA3 silencing led to swollen stems, pedicels, filaments, styles and epidermal hairs as well as thickened leaves and corollas, which were not observed in the *A. thaliana* cev1 mutant, the rsw1 mutant and the antisense AtCESA3 line. Further micrographs showed that PhCESA3 silencing reduced the length and increased the width of cells, suggesting that PhCESA3 silencing inhibits elongation and stimulates radial expansion in petunia.

The morphology of plant organs is highly regulated through cell expansion and cell division1. The final shape and size of cells are often determined by the magnitude and direction of primary cell wall extension2–4. After expansion is arrested, a thick, secondary cell wall is often deposited within the boundary of the primary cell wall5.

In the past decades, significant insight into the molecular details of cellulose biosynthesis has been gained using forward and reverse genetic analyses. The plasma membrane rosettes contain the cellulose synthase catalytic subunit (CESA) proteins6–10. There are multiple CESA genes in the CESA gene family in plant genomes. For example, the *Arabidopsis thaliana* genome contains 10 CESA genes. Among these genes, CESA1, CESA3, and CESA6 are required for cellulose biosynthesis in primary cell walls11, whereas CESA4, CESA7, and CESA8 are required for cellulose biosynthesis during secondary wall deposition6–9.

In *A. thaliana*, the CESA3 mutation (cev1) or antisense lines have altered plant growth, including plant height, leaf size and petal size, as well as altered fertility12–14, which are the important ornamental characteristics for ornamental plants. In addition, CESAs are involved in resistance, activating hormone signaling and cellulose microfibril crystallinity10,12,14–16. Studies on *A. thaliana* and *Nicotiana tabacum* L. variety Samsun NN have demonstrated that the expression of the CESA3 cellulose synthase gene that contains a point mutation, named *ixr1–2*, results in greater conversion of plant-derived cellulose to fermentable sugars17.

*Petunia* (*Petunia hybrida*) is often used as a model for studying ornamental plant growth and development. A developmental map of petunia petals has been assembled by reconstructing the pattern of cell expansion through measurement of cell size18.

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Results

Isolation of *P. hybrida* PhCESA3 cDNA. The full-length cDNA of PhCESA3 was isolated from a cDNA library prepared from the corollas of petunia by homologous cloning and the RACE-PCR method. PhCESA3 was predicted to encode a protein of 1083 amino acids with a calculated molecular weight of 120.8 kDa and a pI of 7.02.

The multiple sequence alignments of PhCESA3 protein in the petunia and *A. thaliana* are presented in Fig. S1. PhCESA3 shares 86.5% amino acid sequence identity with AtCESA3 (Table S1). In addition to a putative zinc-binding domain (at amino acid residues 31–76) and two putative transmembrane helices in the N-terminus, PhCESA3 contains six putative transmembrane helices in the C-terminus (Fig. S1) similar to *A. thaliana* CESA proteins, and the C-terminus is strikingly conserved19,20. In the course of the publication of the study, the petunia genome sequences were released21 (https://www.sgn.cornell.edu/). By blasting against Sol Genomics Network with *A. thaliana* AtCESAs, nine PhCESAs, including PhCESA3, were obtained in the petunia genome. The PhCESAs are named PhCESA1, PhCESA2A, PhCESA2B, PhCESA2C, PhCESA3, PhCESA4, PhCESA6, PhCESA7 and PhCESA8 based on the identity of their encoding amino acid sequences with the *A. thaliana* AtCESAs (Table S1). The phylogenetic tree of the PhCESAs and *A. thaliana* AtCESAs shown in Fig. 1 further reveals that the cloned PhCESA3 is the ortholog of AtCESA3.

Expression patterns of PhCESA3 mRNA. As a step toward functional analysis, we examined the spatiotemporal expression of PhCESA3 petunia 'Ultra' using quantitative real-time PCR (qPCR) with gene-specific primers, and actin was used as an internal control. PhCESA3 transcripts were present in root, stem, leaf, and corolla, with varying patterns of expression (Fig. 2). PhCESA3 mRNA was slightly more abundant in roots and stems compared to corollas.

VIGS-mediated PhCESA3 silencing leads to plant dwarfing. To explore the function of PhCESA3, we generated a TRV-PhCESA3 vector that contains a 266 bp fragment of the 3′ untranslated sequence of PhCESA3 cDNA to test the silencing of PhCESA3 by virus-induced gene silencing (VIGS)22. Five weeks after TRV infection, plants infected with TRV-PhCESA3 showed a significant dwarfing phenotype. The stems and internodes were shorter and wider than those of the empty TRV-infected control plants (Fig. 3a–e; Table 1). The silenced stems were about half the length of the control. The diameter of silenced stems was significantly increased. In addition, there were shorter stem epidermal hairs on PhCESA3-silenced plants compared to control plants (Fig. 3f). PhCESA3 silencing also resulted in smaller and thicker leaves than those of control plants (Fig. 3g and Table 1).

Analysis by qPCR showed decreased PhCESA3 mRNA levels in corollas, leaves and stems in plants treated with TRV-PhCESA3 compared to those in control plants (Figs 4 and S2 and 3).

In addition, PhCESA3 silencing has no significantly effect on the expression of the other 8 PhCESAs in corollas (Fig. 4), leaves (Fig. S2) and stems (Fig. S3). PhCESA2B, PhCESA2C, and PhCESA6 mRNA levels were not detected by qPCR in either TRV-PhCESA3-treated or control corollas and PhCESA6 mRNA levels were also not detected in either TRV-PhCESA3-treated or control leaves.

PhCESA3 silencing alters flower organ growth. PhCESA3 silencing reduced the height of stems but did not change the flowering time. However, altered flower organ growth was observed in PhCESA3-silenced plants. PhCESA3 silencing reduced the size but increased the thickness of petal limbs, and it reduced the height of petal tubes (Fig. 3h and i; Table 1).

The pedicels (fruit stalks), filaments and styles were shorter and wider in PhCESA3-silenced plants than those in control plants (Fig. 3j–l; Table 1). The lengths of pedicels (fruit stalks), styles, and filaments in PhCESA3-silenced plants were about two-thirds, half and half of those of control plants, respectively (Table 1). The diameters of pedicels and styles of PhCESA3-silenced plants were about one and half, and two fold of those of control plants, respectively (Table 1). In addition, the styles and filaments were semi-transparent and loose, and their surface was rough in PhCESA3-silenced plants (Fig. 3n and o). In PhCESA3-silenced plants, the stigma was small and deformed (Fig. 3m), and the unpollinated ovaries and young fruits were often deformed (Fig. 3p and q).

PhCESA3 silencing leads to a significant reduction in cellulose content. To further characterize the function of PhCESA3, cellulose content was analyzed in plants treated with TRV and TRV-PhCESA3 treatment plant. PhCESA3-silenced plants showed a significant decrease in total cellulose content of the stems and leaves (Fig. 5a and b). In 5-week-old plants, mature stems in PhCESA3-silenced plants had about 80% and mature leaves had about 70% of the control level of cellulose (Fig. 5a and b).

PhCESA3 silencing leads to a significant reduction in fertility. The effects of PhCESA3 silencing on fertility were examined. PhCESA3 silencing caused a significant reduction in fertility, resulting in fewer than 100 seeds per fruit. More seeds per fruit were obtained in reciprocal crosses between PhCESA3-silenced plants and control plants when PhCESA3-silenced plants were used as the pollen donor or receptor (Table 2) but fewer than crosses between both control plants as parents, indicating reduced female and male fertility. Reciprocal crosses with control plants showed that the female reproductive capacity was more strongly affected than male reproductive capacity in PhCESA3-silenced plants (Table 2).

PhCESA3 silencing inhibits cell elongation and causes cell swelling. Because PhCESA3 silencing altered the growth of stems, leaves and flowers, we further examined these organs in detail. Scanning electron micrographs showed reduced cell length and increased cell width of the epidermal cells of stems, pedicels, filaments and styles in PhCESA3-silenced plants (Fig. 6a–d; Table 3). Optical micrographs of the epidermis of stems and pedicels showed similar results (Fig. S4a and b).
As visualized in transverse and longitudinal sections, PhCESA3 silencing also reduced the length and increased the width of the cortical cells of stems, pedicels, and filaments, but the number of cortical cells did not change (Fig. 7a,b,e,f and l; Table 3), indicating that the swollen stems were attributed to an increased cell width rather than a decrease in cell number. Hand-cut sections also showed similar results (Fig. S5a and b). The transverse sections showed wider pedicels, filaments and styles in PhCESA3-silenced plants than those in control plants (Fig. 7d,j,m; Table 1). In addition, the arrangement of cortical cells in stems, pedicels and filaments was irregular and close in PhCESA3-silenced plants compared with that of control plants (Fig. 7b,e,l). The cortical cells were often round or oval shaped in control plants, while the cortical cells were often irregular polygon shaped in PhCESA3-silenced plants (Fig. 7b,e and l).

Scanning electron micrographs of adaxial and abaxial epidermal cells of leaves and petal limbs showed reduced cell size in PhCESA3-silenced plants (Fig. 6e and h). The reduced proportion of leaves and petal limbs was similar to that of the epidermal cells of leaves and petal limbs. Optical micrographs of the epidermis of leaves and petals showed similar results (Fig. S4c–f). As visualized in transverse sections, PhCESA3 silencing increased the width

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**Figure 1.** Neighbor-joining trees among proteins encoded by the CESA-like genes using DNAMAN. Petunia PhCESAs were aligned with *Arabidopsis thaliana* AtCESA1 (AT4G32410), AtCESA2 (AT4G39350), AtCESA3 (AT5G05170), AtCESA4 (AT5G44030), AtCESA5 (AT5G09870), AtCESA6 (AT5G64740), AtCESA7 (AT3G17420), AtCESA8 (AT4G18780), AtCESA9 (AT2G21770), and AtCESA10 (AT2G25540). Petunia PhCESAs come from Sol Genomics Network. PhCESA1, Peaxi162Scf00073g01124; PhCESA2A, Peaxi162Scf00025g00276; PhCESA2C, Peaxi162Scf00079g02433; PhCESA2B, Peaxi162Scf01294g00032; PhCESA3, Peaxi162Scf00953g00116; PhCESA4, Peaxi162Scf00415g00078; PhCESA6, Peaxi162Scf00041g00017; PhCESA7, Peaxi162Scf00401g00329; PhCESA8, Peaxi162Scf00371g00318.
of the mesophyll cells of leaves and petals (Fig. 7c and g), while the number of mesophyll cells did not change, indicating that the thickened leaves and petals were attributed to an increased cell size rather than cell number.

In addition, the sections of petal limbs revealed that the outside shape of the epidermal cells of the adaxial side of petal limbs from control plants was often sharp, while the outside shape of the same cells from PhCESA3-silenced plants was curved (Fig. 7g and h). The epidermal cells of the abaxial side of petal limbs of control plants showed a similar size and good arrangement, but the same cells in PhCESA3-silenced plants had different sizes, different shapes, and irregular arrangement (Fig. 7g and i). Transmission electron micrographs of cells from petal limbs confirmed these results (Fig. 8e–g).

Scanning electron micrographs showed that the epidermal cells of petals in PhCESA3-silenced plants were often collapsed while those of control plants were plump (Fig. 6g). The collapsed cells may have been due to thin and fragile epidermal cell walls in PhCESA3-silenced plants compared to those of control plants as revealed by transmission electron microscopy (Fig. 8e).

The transverse sections of filaments and styles showed the wide marrow in PhCESA3-silenced plants compared with that of the control (Fig. 7k and n). Moreover, the myeloid cells of control plants were similar in size and had a good arrangement while those of PhCESA3-silenced plants had different sizes, different shapes and irregular arrangement (Fig. 7k and n). In addition, wider epidermal cells of style were observed in the PhCESA3-silenced plants compared to those in the control plants (Fig. 7o).

Scanning electron and optical micrographs of hand-cut sections and stereomicroscope micrographs all showed that PhCESA3 silencing reduced the length and increased the width of cells of epidermal hairs in stems and pedicels (Fig. 5c and d). Scanning electron microscopy showed the increase of number of epidermal hairs of pedicels in the PhCESA3-silenced plants (Fig. 6i). In addition, scanning electron micrographs showed collapsed epidermal hairs on PhCESA3-silenced plants, which may be attributed to the thin and fragile cell walls in PhCESA3-silenced plants (Fig. 6i).

In addition, scanning electron micrographs of pollen grains showed that parts of the pollen cell wall were collapsed in PhCESA3-silenced plants, but the pollen cells in control plants were plump (Fig. 6k), which may partially explain the sterility of PhCESA3-silenced plants.

**PhCESA3 silencing alters the ultrastructure of cells.** To investigate if PhCESA3 silencing affects the ultrastructure of cells, we examined the thickness of cell walls. Transmission electron micrographs of cell walls revealed that the epidermal cell walls of stems, petal limbs and filaments in PhCESA3-silenced plants were thin compared to those of control plants (Fig. 8a, e, f, g and h; Table 4), while the thickness of the cell walls of cortical cells and xylem cells of stems was not significantly different between the plants (Fig. 8b and d; Table 4). In addition, we observed that the starch granules in chloroplasts of stem cortical cells in PhCESA3-silenced plants were rare compared to control plants (Fig. 8c).

**Discussion**

CESA3, which encodes the catalytic subunit 3 of cellulose synthase, has been shown to be involved in the biosynthesis of primary cell walls in *A. thaliana*. We isolated a petunia homolog of the *A. thaliana* CESA3 gene (PhCESA3), and we analyzed the spatial and temporal regulation of PhCESA3 expression. Furthermore, we produced VIGS-mediated PhCESA3-silenced plants and compared these plants to control plants to examine the role of PhCESA3 in plant growth and development.

The predicted peptide sequence of PhCESA3 shows high similarity to *A. thaliana* CESAs throughout the protein with a putative zinc-binding domain and two putative transmembrane helices in the N-terminus as well as six putative transmembrane helices in the C-terminus (Fig. S1). PhCESA3 has the homology with AtCESA3 in *A. thaliana*, which indicates that PhCESA3 is the homolog of AtCESA3. PhCESA3 mRNA was detected in all organs examined but was present at different levels in different organs (Fig. 2), indicating that PhCESA3 expression is spatially regulated.

In the flowers of plants treated with TRV-PhCESA3, PhCESA3 mRNA levels showed decreased compared to those of control flowers, while the mRNA levels of other PhCESAs were not significantly changed compared to those of control flowers (Fig. 4), which showed the specific silencing of PhCESA3 in plants treated with TRV-PhCESA3.
In *A. thaliana*, mutants of AtCESA4, 7, and 8 specifically show a cellulose defect in the secondary wall of the xylem, whereas AtCESA1, 3 and 6 are mainly involved in cellulose synthesis in the primary cell wall. AtCESA1 and AtCESA3 are required for cellulose synthesis in the primary cell wall, whereas AtCESA2 and AtCESA6 may be at least partially redundant. Null alleles of AtCESA3 and AtCESA1 are embryo lethal. In *A. thaliana*, the rsw1 mutant (the temperature-sensitive allele of AtCESA1) the antisense AtCESA3 line, and the cev1 mutant (a mild mutation of AtCESA3) results in dwarfing of plants, small leaves, small petals, short pedicels, short filaments, short styles, and reduced fertility compared to control plants. The short stamen filament surface of the rsw1 mutant is crumpled. In tobacco, using the VIGS-method silencing of *NtCESA1*, the homolog of...
AtCESA3, leads to shorter internode lengths, small leaves, and a dwarf phenotype\(^30\). In this study, similar results were obtained in the PhCESA3-silenced plants (Fig. 3a–e,g,i–l). Both the antisense AtCESA3 line and rsw1 mutant affect female reproductive competence more severely than male competence in A. thaliana\(^13,29\), which is consistent with the results in this study (Table 2). However, PhCESA3 silencing caused swollen stems, pedicels, filaments, styles and epidermal hairs, which were not observed in the AtCESA3 antisense line, the cev1 mutant and the rsw1 mutant in A. thaliana\(^12–15,29\). Further, scanning, transmission and optical micrographs showed that PhCESA3 silencing caused short and swollen cells of these organs or tissues, which was consistent with the corresponding phenotypes. Thus, PhCESA3 silencing resulted in primary cell wall extension in length and width. Importantly, growing the rsw1 mutant at its restrictive temperature inhibits root and hypocotyl elongation as well as promotes radial swelling\(^29\). Mutations in A. thaliana AtCESA3 and AtCESA6 confer resistance to isoxaben (an herbicide that

|                     | TRV (Control) | TRV-PhCESA3 | TRV-PhCESA3/TRV (%) |
|---------------------|---------------|-------------|---------------------|
| Height of plant (cm) | 18.75 ± 1.28  | 10.65 ± 2.03* | 57                  |
| Length of internode (cm) | 2.8 ± 0.58  | 0.68 ± 0.20*  | 24                  |
| Diameter of stem (mm)  | 2.25 ± 0.21  | 3.73 ± 0.52*  | 166                 |
| Length of leaf (cm)    | 4.45 ± 0.58  | 3.32 ± 0.23*  | 75                  |
| Width of leaf (cm)     | 2.46 ± 0.31  | 1.84 ± 0.26*  | 75                  |
| Thickness of leaf (µm) | 198.67 ± 32.55 | 337.93 ± 65.96* | 170                |
| Length of pedicel (cm) | 3.88 ± 0.9   | 1.15 ± 0.28*  | 30                  |
| Diameter of pedicel (mm) | 1.82 ± 0.21  | 2.75 ± 0.36*  | 151                 |
| Diameter of corolla (cm) | 6.71 ± 0.48  | 3.81 ± 0.65*  | 57                  |
| Thickness of petal limb (µm) | 103.24 ± 12.91 | 160.92 ± 33.81* | 156                |
| Length of filament (cm) | 1.19 ± 0.54  | 0.62 ± 0.17*  | 52                  |
| Diameter of filament (mm) | 0.44 ± 0.03  | 0.80 ± 0.17*  | 182                 |
| Length of style (cm)   | 2.16 ± 0.08  | 1.30 ± 0.21*  | 60                  |
| Diameter of style (mm) | 0.62 ± 0.07  | 1.20 ± 0.24*  | 194                 |
| Thickness of stem cortex (µm) | 174.72 ± 24.43 | 301.69 ± 37.68* | 173                |
| Thickness of pedicel cortex (µm) | 277.22 ± 33.74 | 359.73 ± 60.44* | 130                |

Table 1. PhCESA3 silencing alters plant growth. Data are means ± SE from 15 to 20 samples. Statistical analysis was performed using Student t test with 15 to 20 replicates. Asterisk means significant difference at \(P = 0.05\) level.

Figure 4. Effects of TRV2-PhCESA3 treatment on the expression of PhCESA3 and other PhCESAs in flowers on day 2 after opening as determined by quantitative real-time PCR. Relative expression levels are shown as fold change values. Data are presented as the mean ± SD \((n = 3)\). Statistical analysis was performed using Student t test with 3 replicates. Asterisk means significant difference at \(P = 0.05\) level.
specifically inhibits cellulose) and isoxaben, which causes radial swelling of roots. Taken together, with strong inhibition of cellulose synthesis, whether genetic or chemical (i.e., with herbicides such as isoxaben), organ elongation is inhibited and radial expansion is stimulated. However, the organs that exhibit radial expansion induced by \( \text{CESA} \) silencing are different in various species.

Among the ten \( \text{AtCESAs} \) in \( A. \text{thaliana} \), \( \text{AtCESA3} \) (\( \text{IXR1/CEV1} \)) is suggested to be essential for depositing cellulose in primary walls. When changes to primary walls alter cell expansion and/or cell divisions, changes to cell shape, growth, and morphogenesis occur. Changes to thickening after growth stops occur too late to affect these processes, resulting in normal morphology but changes in mechanical properties. Similar to the \( A. \text{thaliana} \) \( \text{AtCESA3} \) mutant, \( \text{PhCESA3} \) silencing resulted in significant changes in the cell size and shape in several organs. Moreover, the cellulose content of mature stems and leaves from \( \text{PhCESA3} \)-silenced plants was significantly decreased compared to that of control plants. These results suggested that \( \text{PhCESA3} \) mainly functions for catalyzing the biosynthesis of cellulose deposited to the primary wall. Ultrastructural studies clearly showed that the cell walls of the epidermis in stems and petals, which are heavily thickened during primary growth, were thinner, thus further supporting the involvement of \( \text{PhCESA3} \) in depositing cellulose in the primary wall. The thickness of the cell walls of cortical cells and xylem cells of stems did not significantly change, suggesting that \( \text{PhCESA3} \) is not involved in depositing cellulose in thickening walls after cell growth stops.

In this study, the shorter stems, pedicels, filaments and styles in \( \text{PhCESA3} \)-silenced plants were attributed to reduction in cell length rather than cell number, and these swollen organs in \( \text{PhCESA3} \)-silenced plants were attributed to increased cell width rather than cell number. These results implicated that \( \text{PhCESA3} \) is mainly involved in cell elongation rather than cell division, which is consistent with results obtained with the antisense \( \text{AtCESA3} \) line in \( A. \text{thaliana} \). In addition, plant organ shaping requires control over cell wall expansion anisotropy, which is characterized by the direction and degree of anisotropy. In this study, the inhibited elongation and stimulated radial expansion of several organs in \( \text{PhCESA3} \)-silenced plants showed the change of the direction and degree of anisotropy of cell wall.

Here, \( \text{PhCESA3} \) silencing resulted in changes of plant height, flower size and fertility, which are the important features of ornamental plants. These results suggested the important role of \( \text{PhCESA3} \) in controlling plant stature and form. By using specific promoters, changes in \( \text{CESA3} \) expression in different organs may create an excellent variety for ornamental plants. Therefore, \( \text{PhCESA3} \) may be utilized in petunia breeding for changing the plant height, flower size and fertility.

![Cellulose content of mature stems and leaves from wild-type and \( \text{PhCESA3} \)-silenced plants.](image)

**Figure 5. Cellulose content of mature stems and leaves from wild-type and \( \text{PhCESA3} \)-silenced plants.** Cellulose was measured in stem segments (a) and leaves (b) from wild-type and \( \text{PhCESA3} \)-silenced plants. Standard error bars are shown (\( n = 6 \)). Statistical analysis was performed using Student t test with 6 replicates. Asterisk means significant difference at \( P = 0.05 \) level.

|                      | TRV self-fertilization | TRV-\( \text{PhCESA3} \) self-fertilization | TRV \( \varphi \) and TRV-\( \text{PhCESA3}\) \( \varphi \) | TRV-\( \text{PhCESA3}\) \( \varphi \) and TRV \( \delta \) |
|----------------------|------------------------|--------------------------------------------|------------------------------------------------|------------------------------------------------|
| Number of seeds per fruit | 205 ± 13a              | 92 ± 11d                                   | 155 ± 12b                                       | 118 ± 12c                                       |

**Table 2. Effects of \( \text{PhCESA3} \) silencing on the fertility.** Data are means ± SE from 15 to 20 fruits. Different letters mean significant difference at \( P = 0.05 \) level. Statistical analysis was performed using one way analysis of variance (ANOVA) followed by Duncan’s multiple range test (DMRT) with 15 to 20 replicates. \( P \)-values \( \leq 0.05 \) were considered as significant.
Methods

Plant material. Petunia 'Ultra' plants were grown under greenhouse conditions (22–25 °C, 14 h light/10 h dark). Flowers were emasculated 1 d before the flowers were fully open to prevent self-pollination. Eight to ten petunia flowers were harvested at anthesis (corollas 90° reflexed) stages and were then placed immediately in tap water. Stem, leaves and roots were collected from plants at the vegetative stage when the plants were approximately 10 cm in height. All tissues were frozen in liquid nitrogen and stored at −80 °C until used for RNA extraction. Fresh weights were measured immediately before freezing. All experiments were conducted at least three times with independently collected and extracted tissues unless otherwise noted.

RNA extraction, RT-PCR and cloning of the petunia PhCESA3 gene. Total RNA was extracted and reverse-transcribed according to the methods of Liu et al. The partial sequences of PhCESA3 were obtained using a computational identification approach. Briefly, TBLASTN analysis against the GenBank EST database (http://www.ncbi.nlm.nih.gov) with A. thaliana CESA3 identified 1 petunia clone, FN016492, which encodes putative proteins that displayed conservation with A. thaliana CESA3. The remaining 5' and 3' cDNA sequences of PhCESA3 were isolated by RACE using a specific primer (Table S2), and its full-length cDNA was isolated by RT-PCR.

Figure 6. Scanning electron micrographs of the PhCESA3-silenced plants compared with those of control plants. (a–d), Epidermal cells of stem (a), pedicel (b), filament (c) and style (d) showing reduced cell length and increased cell width in PhCESA3-silenced plants (right, bottom in (d)) compared to control plants (left, top in (d)). (e,f), Adaxial (e) and abaxial (f) epidermal cells of leaves showing the reduced cell size in PhCESA3-silenced plants (right) compared to control plants (left). (g,h), Adaxial (g) and abaxial (h) epidermal cells of petal limb showing the reduced cell size in PhCESA3-silenced plants (right) compared to control plants (left). (i), Pedicels showing the reduced number of trichomes in PhCESA3-silenced plants (right) compared to control plants (left). (j) Trichomes of pedicel showing the reduced trichome length and more collapsed trichomes in PhCESA3-silenced plants (right) compared to control plants (left). (k), Pollen grains showing the abnormal pollen mother cell in PhCESA3-silenced plants (right) compared to control plants (left). Bars = 50 μm in (a–f); bars = 20 μm in (g–h); bar = 1 mm in (i); bars = 200 μm in (j); bars = 10 μm in (k).
10 kV acceleration and photographed. Samples were observed with a scanning electron microscope (XL-30-ESEM, The Netherlands, FEI) at Switzerland, Bal-Tec). The dried samples were fixed on the sample stage and coated with gold by ion sputtering. Samples were dehydrated in increasing grades of ethanol and then dried with a critical point drier (CPD 030, Glutaraldehyde in 0.1 mol/L PBS (pH 7.2) for 4 h at 4°C and then washed 3 times in the same buffer, which was followed by post-fixation in 1% osmium tetroxide for 2 h at room temperature and 3 rinses using the same buffer. Analyses were conducted following the Minimum Information for Publication of Quantitative Real-Time PCR Experiments guidelines. Three Agrobacterium (strain GV3101) transformed with pTRV1 and pTRV2 derivatives were prepared as previously described. The Agrobacterium cells grown overnight were harvested and resuspended in inoculation buffer containing 10 mM MES, 200 mM acetosyringone, and 10 mM MgCl2 to an OD550 of 10. Following an additional 3 h of incubation, 200 to 400 μL of this mixture was injected into the stem or applied on the cut surface after removing the apical meristems of petunia plantlets. Approximately thirty plants were vaccinated with each vector. The inoculated plants were grown under greenhouse conditions (22–25 °C, 14 h light/10 h dark).

**Table 3. Effects of PhCESA3 silencing on the cell size** Data are means ± SE from 15 to 20 samples. Statistical analysis was performed using Student t test with 15 to 20 replicates. Asterisk means significant difference at P = 0.05 level.

|                          | TRV (Control) | TRV-PhCESA3 | TRV-PhCESA3/TRV (%) |
|--------------------------|--------------|-------------|---------------------|
| Height of epidermal cells of stem (μm) | 89.69 ± 14.17 | 50.74 ± 14.59* | 57                  |
| Width of epidermal cells of stem (μm)    | 28.50 ± 5.56  | 55.47 ± 14.29* | 209                 |
| Height of cortex cells of stem (μm)      | 117.07 ± 19.06 | 67.83 ± 16.42* | 58                  |
| Diameter of adaxial epidermal cells of leaf (μm) | 101.52 ± 6.23 | 84.59 ± 7.86*  | 83                  |
| Height of cortical cells of leaf (μm)    | 93.39 ± 5.67  | 76.67 ± 6.46*  | 82                  |
| Height of epidermal cells of pedicel (μm) | 166.09 ± 56.89 | 66.76 ± 18.63* | 40                  |
| Diameter of abaxial epidermal cells of leaf (μm) | 161.02 ± 20.30 | 83.41 ± 33.94* | 52                  |
| Height of cortical cells of pedicel (μm) | 58.76 ± 9.38  | 117.77 ± 20.56* | 200                 |
| Height of cortical cells of filament (μm) | 65.33 ± 7.67  | 25.78 ± 4.60*  | 41                  |
| Width of epidermal cells of filament (μm) | 18.88 ± 3.20  | 30.99 ± 6.02*  | 164                 |
| Height of epidermal cells of style (μm)  | 29.46 ± 2.84  | 76.94 ± 14.44* | 261                 |
| Height of epidermal cells of stamen (μm) | 119.07 ± 14.13 | 62.74 ± 14.91* | 53                  |
| Width of epidermal cells of stamen (μm)  | 16.78 ± 2.81  | 26.60 ± 5.62*  | 159                 |

**Sequence analysis.** Alignments were performed, and a phylogenetic tree was generated using the DNAMAN software. Identity search for nucleotides and translated amino acids was performed using the National Center for Biotechnology Information (NCBI) BLAST network server (http://www.ncbi.nlm.gov/BLAST).

**Quantitative real-time PCR assays.** Quantitative real-time PCR (qPCR) assays were performed according to previous methods. Analyses were conducted following the Minimum Information for Publication of Quantitative Real-Time PCR Experiments guidelines. Petunia Actin (accession no. FN014209) and Cyclophilin (CYP) (accession no. EST883944) genes were used as the internal reference genes to quantify the cDNA abundance. Similar results were obtained for both reference genes. The data presented in the body of the text represent relative expression values calculated using Actin, and the data presented in the attachment represent relative expression values calculated using CYP. The sequences of all primers used for qPCR analysis are described in Table S3.

**Agroinoculation of TRV vectors.** To generate pTRV2 containing the 3′ untranslated region of PhCESA3 (TRV2-PhCESA3), the gene sequence of 266 bp was PCR amplified using forward primers and reverse primers (Table S4), and the PCR products were inserted into the pTRV2 vector. Agrobacterium tumefaciens (strain GV3101) transformed with pTRV1 and pTRV2 derivatives were prepared as previously described. The Agrobacterium cells grown overnight were harvested and resuspended in inoculation buffer containing 10 mM MES, 200 mM acetosyringone, and 10 mM MgCl2 to an OD550 of 10. Following an additional 3 h of incubation at 28°C, bacteria transformed with pTRV1 were mixed with bacteria containing the pTRV2 derivatives in a 1:1 ratio, and 200 to 400 μL of this mixture was injected into the stem or applied on the cut surface after removing the apical meristems of petunia plantlets. Approximately thirty plants were vaccinated with each vector. The inoculated plants were grown under greenhouse conditions (22–25°C, 14 h light/10 h dark).

**Cellulose measurement.** Cellulose content in mature stems and leaves was measured by a previously published method. Analysis was conducted with material from a single plant using three to five fully expanded leaves from 3-week-old plants or a 3 cm segment from the base of the primary inflorescence stem. Stems were chopped with a razor blade, and a crude cell wall fraction was obtained by extracting the soluble material with two changes of 70% ethanol at 70°C for 1 h each. The ethanol was removed, and the samples were dried under vacuum. The dry weight of the wall material was recorded. Cellulose was measured according to a previously published method.

**Scanning electron micrographs.** The stems, leaves, pedicels, petal limbs, filaments, styles and anthers from PhCESA3-silenced plants and control plants were cut into 3–5 mm2 pieces. The samples were fixed in 4% glutaraldehyde in 0.1 mol/L PBS (pH 7.2) for 4 h at 4°C and then washed 3 times in the same buffer, which was followed by post-fixation in 1% osmium tetroxide for 2 h at room temperature and 3 rinses using the same buffer. Samples were dehydrated in increasing grades of ethanol and then dried with a critical point drier (CPD 030, Switzerland, Bal-Tec). The dried samples were fixed on the sample stage and coated with gold by ion sputtering equipment. Samples were observed with a scanning electron microscope (XL-30-ESEM, The Netherlands, FEI) at 10 kV acceleration and photographed.
The stems, leaves, pedicels and styles from plants were cut into 5 mm × 5 mm × 5 mm pieces. The samples were fixed in FAA fixative solution (every 100 ml of FAA fixative solution contains 90 ml of 50% or 70% ethanol, 5 ml of acetic acid, and 5 ml formalin) for 24 h at room temperature and then washed in running water for 24 h. Samples were stained by hematoxylin for 4 d and then washed in running water for 24 h, which was followed by dehydration in increasing grades of ethanol. A graded chloroform series was used for clearing, and the samples were embedded in paraffin. Paraffin sections were cut to a thickness of 8 μm on a Leica RM2235 followed by dewaxing with xylene. Lastly, slides were sealed with neutral resin. Sections were observed and photographed with a Zeiss Scope.A1 microscope.

**Figure 7.** Section micrographs of PhCESA3-silenced plants compared to control plants. (a) Stem longitudinal section showing the increased width of cortex and cortical cells in PhCESA3-silenced plants (right) compared to control plants (left). (b) Stem transverse section showing the uneven surface in PhCESA3-silenced plants (right) compared to control plants (left). (c) Leaf transverse section showing the thickened leaf in the PhCESA3-silenced plants (right) compared to the control plant (left). (d) Pedicel transverse section showing the thickened pedicel in the PhCESA3-silenced plants (right) compared to the control plant (left). (e,f) Pedicel transverse (e) and longitudinal (f) sections showing the increased width of cortical cells in PhCESA3-silenced plants (right) compared to control plants (left). (g) Petal limb transverse section showing the thickened petal limb in the PhCESA3-silenced plants (right) compared to the control plant (left). (h) Solid and hollow arrows indicate the cell shape of the adaxial and abaxial sides, respectively. (i) The adaxial epidermal cells of petal limb showing sleek cells in the PhCESA3-silenced plants (bottom) compared to the control plant (top). (j) The abaxial epidermal cells of petal limb showing the uneven size and irregular shape of cells in the PhCESA3-silenced plants (bottom) compared to the control plant (top). (j,m) Transverse section showing the thickened filament (j) and style (m) in the PhCESA3-silenced plants (right) compared to the control plant (left). (k,n) Marrow transverse section of filament and style showing the irregularly arranged marrow cells and different cell sizes in the PhCESA3-silenced plants (right) compared to the control plant (left). (l) Filament cortex transverse section showing the irregularly arranged cortical cells and uneven cell sizes in the PhCESA3-silenced plants (right) compared to the control plant (left). (o) Style transverse section showing the width of epidermal cells in the PhCESA3-silenced plants (right) compared to the wild-type plant (left). The cortex (co), cambium (ca), and xylem elements (xe) are indicated. Bars = 250 μm in (a,b,f,j,m); bars = 100 μm in (e,c); bars = 500 μm in (d,l); bars = 50 μm in (g,k,n,o); bars = 10 μm in (h,i).
Semi-thin sections. The pedicels, petal limbs and filaments from PhCESA3-silenced plants and control plants were cut into 1 mm × 1 mm × 0.5 mm pieces. The specimens were fixed in 2.5% paraformaldehyde/3.0% glutaraldehyde in 0.1 mol/L PBS (pH 7.2) for 4 h at 4 °C, and the specimens were then washed 3 times in the same buffer, which was followed by post-fixation in 1% osmium tetroxide for 2 h at room temperature and 3 rinses using the same buffer. The specimens were dehydrated in a graded ethanol series and embedded in Epon812 (SPI...
Supplies Division of Structure Probe Inc., West Chester, PA, USA). Polymerization took place for 24 h at 40 °C, which was followed by 24 h at 60 °C. Specimens were cut to a thickness of 1 μm on a Leica RM2155 and were stained with 0.5% toluidine blue. Sections were observed and photographed with a Leica DMLB microscope.

Transmission electron micrograph. The stems, petal limbs and filaments from PhCESA3-silenced plants and control plants were cut into <1 mm³ pieces. The samples were fixed in 4% glutaraldehyde in 0.1 mol/L PBS (pH 7.2) for 4 h at 4 °C and then washed 3 times in the same buffer, which was followed by post-fixation in 1% osmium tetroxide for 2 h at room temperature and 3 rinses using the same buffer. Samples were dehydrated in increasing grades of ethanol and embedded in Epon812 (containing 51.64% Epon812, 5.37% DDSA, 42.99% MNA and 1.5% DMP-30). Polymerization took place for 24 h at 45 °C, which was followed by 24 h at 60 °C. Ultrathin sections (100 nm thick) were cut with a Leica ULTACUT ultramicrotome and deposited on a 200 mesh copper net with support film. Samples were stained with a 2% uranyl acetate solution followed by a 6% lead citrate solution, and samples were visualized with a Tecnai 12 transmission electron microscope (FEI, Eindhoven, The Netherlands) at 80 kV acceleration and photographed.

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**Author Contributions**

Yu Y. and Liu J. designed the research. Yang W., Cai Y., Hu L., Wei Q., Chen G., Bai M., and Wu H. performed the research. Yu Y. and Yang W. wrote the paper.

**Additional Information**

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