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


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Inflorescence abscission protein SIIDL6 promotes low light intensity-induced tomato flower abscission

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Abstract

In many fruiting plant species, flower abscission is induced by low light stress. Here, we elucidated how signaling mediated by the peptide INFLORESCENCE DEFICIENT IN ABSCISSION (IDA) controls low light-induced flower drop in tomato (*Solanum lycopersicum*). We analyzed the expression patterns of an IDA-Like gene (*SIIDL6*) during low light-induced flower abscission, and used tandem mass spectrometry to identify and characterize the mature SIIDL6 peptide. Tomato knockout lines were created to investigate the in vivo function of SIIDL6. In addition, yeast one-hybrid assays were used to investigate the binding of the SIWRKY17 transcription factor to the *SIIDL6* promoter, and silencing of *SIWRKY17* expression delayed low light-induced flower abscission. *SIIDL6* was specifically expressed in the abscission zone and at high levels during low light-induced abscission and ethylene treatment. *SIIDL6* knockout lines showed delayed low light-induced flower drop, and the application of SIIDL6 peptide accelerated abscission. Overexpression of *SIIDL6* rescued the *ida* mutant phenotype in *Arabidopsis* (*Arabidopsis thaliana*), suggesting functional conservation between species. SIIDL6-mediated abscission was via an ethylene-independent pathway. We report a SIWRKY17-SIIDL6 regulatory module that functions in low light promoted abscission by increasing the expression of enzymes involved in cell wall remodeling and disassembly.

Introduction

Plants have the capacity to shed their vegetative and reproductive organs in response to developmental signals and environmental stimuli (Addicott, 1982). Although this abscission process has many adaptive benefits, in an agricultural context,

premature abscission can cause a major reduction in crop productivity, as exemplified by fruit set failure. Thus, there is considerable interest in understanding the mechanism underlying organ abscission and its regulation.

Abscission is a highly regulated process involving structural, biochemical, and molecular changes in a specialized

tissue of the organ called the abscission zone (AZ; Sexton and Roberts, 1982; Estornell et al., 2013). These changes include protein degradation, cell wall remodeling, dissolution of the middle lamella, loss of cell membrane permeability, and programmed cell death (Cho et al., 2008; Meir et al., 2010; Bar-Dror et al., 2011).

Multiple phytohormones are known to play roles in triggering and modulating abscission, and ethylene is a well-documented accelerator of leaf, flower, and fruit abscission. Accordingly, enhancing ethylene production and promoting ethylene signaling by elevating the expression of ethylene response genes, including transcription factor (TF) genes, have been shown to promote abscission (Butenko et al., 2006; Wilson et al., 2011). In tomato (*Solanum lycopersicum*), transcriptome analysis of the AZ pedicel indicated that ethylene biosynthesis and related signal transduction, including the expression of bZIP and WRKY TF family members, is upregulated during abscission, and pretreatment with the ethylene inhibitor 1-methylcyclopropene (1-MCP) suppressed this upregulation (Meir et al., 2010). Furthermore, WRKY genes were also identified as abscission-specific transcriptional regulators of Arabidopsis (*Arabidopsis thaliana*) flower abscission and soybean (*Glycine max*) leaf abscission (Nath et al., 2007; Niederhuth et al., 2013). However, while WRKY TFs are known to integrate ethylene responses during senescence, their role in abscission is not well understood (Robatzek and Somssich, 2001; Koyama, 2014).

In addition to ethylene, a small peptide, INFLORESCENCE DEFICIENT IN ABSCISSION (IDA), has been identified as an essential signaling element in the induction of Arabidopsis floral organ abscission (Butenko et al., 2003; Stenvik et al., 2008). This IDA signal functions during phases 3 to 4 of *A. thaliana* abscission, which are the last steps, and it has also been shown that the *ida* mutant was completely blocked during flower abscission (Butenko et al., 2003). Before cleavage, IDA-LIKE (IDL) and IDA peptides are usually smaller than 100 amino acids and consist of an N-terminal export signal, a variable region, and a C-terminal extension with a PIP motif. Overexpression of AtIDL1-5 compensated for the abscission defect in the *ida* mutant (Stenvik et al., 2008). Using both the AtIDA promoter to drive the litchi (*Litchi chinensis*) *LcIDL1* gene and the constitutive 35S promoter to drive the *Citrus* IDA3 gene resulted in rescue of the abscission-deficient phenotype of the *ida* mutant, suggesting functional conservation of IDA peptides between species (Estornell et al., 2015; Ying et al., 2016). Five tomato IDA homologs have also been shown to have specific expression patterns during pedicel abscission (Tucker and Yang, 2012). In vitro analysis indicated that the mature *A. thaliana* IDA peptide is 14 amino acids long after cleavage between a Lys and a Gly residue by the subtilase (SBT) family proteases SBT4.12, SBT4.13, or SBT5.2 (Schardon et al., 2016). The leucine-rich repeat (LRR) receptor-like kinases (RLK) HAESA (HAE) and HAESA-LIKE2 (HSL2) are thought to act as IDA receptors. Indeed, the *hae hsl2* double mutant has been reported to have a similar phenotype to the *ida* mutant

(Cho et al., 2008; Stenvik et al., 2008). Other studies have indicated that the hydroxylated IDA peptide binds to HAE and HSL2, and a structural model of the ligand and receptor has been proposed (Santiago et al., 2016). In *A. thaliana*, the KNOTTED-LIKE FROM ARABIDOPSIS THALIANA (KNAT) TFs and cell wall modifying proteins have been identified as downstream components in the IDA-HAE/HSL2 pathway (Shi et al., 2011).

An essential step in abscission is the loss of cell–cell adhesion, which is associated with degradation of the pectin-rich middle lamella. Consistent with this idea, disruption of the expression of an abscission-related pectinase, polygalacturonase, in *A. thaliana* has been shown to significantly delay floral organ drop (González-Carranza et al., 2007). In tomato, abscission-related polygalacturonases (TAPG 1, -2, -3, and -4) are specifically expressed in the flower pedicel AZ during abscission, and their silencing significantly delayed leaf abscission (Jiang et al., 2008). Furthermore, endo- β -1,4-glucanases Cel1 and Cel2, which have been associated with cell wall modification, are expressed in ripening fruit and abscising flowers, and anti-sense based silencing of Cel1 has been shown to delay tomato flower abscission (Gonzalez-Bosch et al., 1997; Lashbrook et al., 1998). In *A. thaliana*, IDA signaling has been reported to be independent from ethylene-triggered abscission, although its transcript levels are lower in *etr1-1* lines (Butenko et al., 2006). However, the role of IDA signaling in tomato abscission and its relationship with ethylene is still unclear.

Drought and low light stress can trigger flower, flower bud, leaf, and fruit abscission in many plant species. In pepper (*Capsicum annuum*), 80% shade for 6 d was shown to significantly promote the abscission of flower buds and positively corresponded with ethylene levels (Wien et al., 1996). However, application of the synthetic auxin 1-naphthaleneacetic acid (NAA) effectively prevented low light-induced flower bud abscission. Furthermore, an eight-fold increase in ethylene production was observed in dark-induced lily (*Lilium candidum*) flower buds during abscission, and application of the ethylene inhibitor silver thiosulfate abolished dark-induced abscission (Van Meeteren and De Proft, 1982). The expression of IDA and IDL genes is also induced by biotic and abiotic stresses (Vie et al., 2015; Patharkar and Walker, 2016). Although IDA-HAE/HSL2 is known to be involved in drought-induced leaf abscission, it is not known whether IDA signaling also mediates low light stress-induced abscission.

Recently, the phyto-sulfokine (PSK; Tyr[SO₃H]-Ile-Tyr[SO₃H]-Thr-Gln) signal has been reported to function in drought-induced tomato flower abscission by promoting the expression of the abscission-related polygalacturonase TAPG4 in an auxin- and ethylene-independent manner (Reichardt et al., 2020). In tomato, PSK is perceived by membrane-bound PSK receptors (PSKR1 and PSKR2), which triggers downstream signaling (Zhang et al., 2018). Thus, PSK and IDA are both involved in abscission and have been associated with cell wall remodeling, but act through different

receptors, and their relationship and relative contributions to the abscission process are unclear.

In this study, we investigated the regulatory mechanism underlying low light-induced flower abscission in tomato, one of the most economically important vegetable crops. Blossom drop is a common phenotype in tomatoes and peppers suffering from dark and low light stress. We report that a low light-induced tomato IDA-like gene, *SIIDL6*, is involved in abscission.

Results

Expression of *SIIDL6* is induced in the AZ following low light-induced abscission

We identified 11 IDA-like genes, including the previously identified *SIIDL1-5* (Tucker and Yang, 2012), using a TBLASTN search (<https://blast.ncbi.nlm.nih.gov/>) with *AtIDA* and *AtIDL* sequences from the tomato genome sequence (Supplemental Figure S1). We named them *SIIDA* and *SIIDL1-10* (Supplemental Table S2). To establish the relationship between *SIIDA* family genes and low light-induced abscission, the expression patterns of *SIIDA/SIIDL* genes were evaluated in a transcriptome database of low-light-treated pedicels. This showed a significant upregulation of *SIIDL6* by the treatment compared with the control conditions (Supplemental Table S3). A RT-qPCR analysis further revealed that *SIIDL6* was specifically expressed in the AZ with a low expression in the distal and proximal regions (Figure 1A), and expression was induced in the AZ under low light stress (Figure 1B). *SIIDL6* expression was significantly increased in the AZ after flower removal induced abscission (Figure 1C). Pedicels treated with ethylene or an ethylene inhibitor (1-MCP) showed elevated or suppressed *SIIDL6* expression, respectively (Figure 1D). By in situ hybridization, we observed that *SIIDL6* was specifically expressed in the AZ and a particularly strong signal was present in the vascular tissue, while no signal was found in the distal or proximal regions, which was consistent with the RT-qPCR results (Figure 1, E–G).

As further evidence that the *SIIDL6* peptide is involved in abscission, the mature 12-mer *SIIDL6* peptide was detected by liquid chromatography mass spectrometry (LC–MS/MS) in culture exudates from the pedicel AZ (Figure 1H; Supplemental Table S4).

Flower drop is enhanced under low light stress in a *SIIDL6* dependent manner

We generated *SIIDL6* knockout lines by CRISPR/Cas9 to investigate the possible role of *SIIDL6* signaling during low light stress-induced flower abscission. Eight independent knockout transgenic lines were obtained through *A. tumefaciens*-mediated transformation (Supplemental Figure S2), five of which were genome deleted. Three independent Cas9-free T2 plants carrying mutations were screened and identified (CR-). In addition, off-target analysis suggested that no off-target activity occurred in the *CRslid6* lines (Table S5). Delayed flower abscission was observed in the knockout

lines under low light stress, and the flower drop was $8.93\% \pm 0.82\%$ compared with $26\%.3 \pm 2.28\%$ in the wild-type (Figure 2, A and B). We further analyzed the effect of *CRslid6* on flower removal abscission in explants. The wild-type plants showed 50% abscission at 16 h and 100% at 40 h, while the *CRslid6* knockout plants had significantly delayed abscission, with pedicels that showed only 50% abscission at 40 h (Figure 2C). To investigate the effect of the *SIIDL6* peptide on pedicel abscission, wild-type and *CRslid6* pedicels were incubated in a medium containing various concentrations of the 12 aa synthetic *SIIDL6* peptide. A *SIIDL6* peptide of 1- μ M had little effect on wild-type abscission, but the 5- and 10- μ M *SIIDL6* peptides significantly enhanced abscission. Notably, 5- and 10- μ M *SIIDL6* peptides had similar effects on abscission, suggesting that the amount of active peptide was saturated at 5- μ M concentration. Furthermore, 5- μ M *SIIDL6* peptide also complemented the *CRslid6* abscission defect (Figure 2D), indicating that tomato flower abscission is regulated by *SIIDL6*.

35S::*SIIDL6* can rescue the *ida2* abscission defect

To test whether *SIIDL6* has a similar role to *AtIDA* in control of floral organ abscission, we generated a construct where *SIIDL6* expression was driven by the cauliflower mosaic virus 35S promoter and transformed into *A. thaliana ida2* mutant plants. Five out of 20 plants were shown to have been transformed, and three lines with high expression were chosen for further experiments (Supplemental Figure S3A). The *ida2* mutants had a flower organ abscission defect phenotype at anthesis, while the *ida2 35S::SIIDL6* transgenic plants showed wild-type phenotype (Supplemental Figure S3B). Furthermore, measurement of the force of petal attachment by petal break strength (pBS) also revealed that overexpression of *SIIDL6* was sufficient to complement the abscission defect in *ida2* (Supplemental Figure S4).

SIIDL6 induced abscission via a partially ethylene-independent manner

To understand whether *SIIDL6* regulates abscission through ethylene signaling, *CRslid6* and wild-type abscission assays were performed following ethylene treatment. In contrast to the wild-type, abscission was observed to be significantly delayed, but not entirely blocked in the *CRslid6* lines, while application of 5- μ M *SIIDL6* peptide accelerated abscission (Figure 3A). We also used the ethylene inhibitor 1-MCP to pretreat the wild-type explants before incubating them with 5- μ M *SIIDL6* peptide, while the controls were not treated. Abscission was significantly accelerated in peptide-treated pedicels compared with the control (Figure 3B). Furthermore, application of 5- μ M *SIIDL6* peptide to the tomato ethylene response mutant, *Neverripe* (Nr), led to faster abscission (Figure 3C). However, it is notable, 5- μ M *SIIDL6* could not restore the abscission of Nr- and 1-MCP-treated plants to levels of the wild-type control. These results indicate that *SIIDL6* can induce abscission via both ethylene-dependent and -independent pathway.

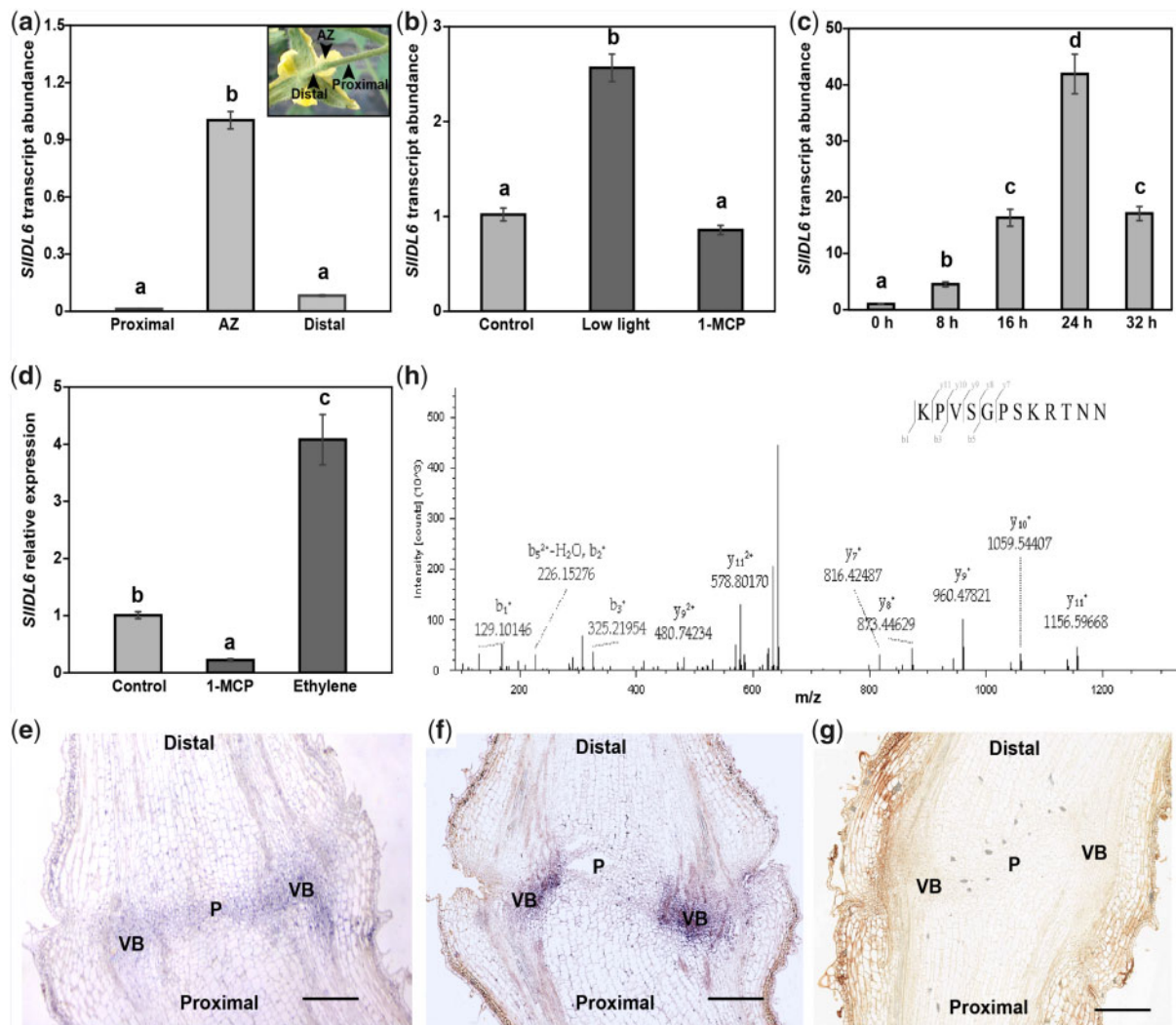


Figure 1 *SIIDL6* expression analysis in flower pedicels. A, The distal, proximal, and abscission zones (AZ) of flower pedicels were used to study *SIIDL6* expression by quantitative real-time PCR. B, *SIIDL6* expression levels in the pedicel AZ under low light treatment compared with the control (normal light), and low light-induced *SIIDL6* expression with 1-methylcyclopropane (1-MCP) treatment. C, The effect of flower removal on *SIIDL6* expression patterns in the wild-type pedicel AZ. D, The effect of $10 \mu\text{L}\cdot\text{L}^{-1}$ ethylene and $2 \mu\text{L}\cdot\text{L}^{-1}$ 1-MCP on *SIIDL6* expression in the pedicel AZ. E–G, Expression analysis of *SIIDL6* at different development stages of pedicel abscission by in situ hybridization. Signal in the pedicel AZ at 8 h (E) or 16 h (F) after flower removal, and in a negative control stained with the positive probe (G). Scale bar = 100 μm . H, LC–MS/MS revealing 12 amino acid mature *SIIDL6* peptide sequences in the tomato pedicel AZ. Different letters indicate significant differences (Student's *t* test, $P < 0.05$). The results are the means of three replicates \pm SD. P, pith; VB, vascular bundle.

Delayed abscission caused by knocking out *SIIDL6* is associated with cell wall degradation

We observed that the expression levels of genes encoding cell wall degrading proteins associated with abscission, including *TAPG1* (Solyc02g067630.2), *TAPG4* (Solyc12g096750), and *CEL2* (Solyc09g010210.2), were downregulated in *CRslid6* lines compared with the wild-type and upregulated by the peptide treatment (Figure 4, A–C). In addition, $P_{TAPG4}::GUS$ staining intensity was enhanced by the *SIIDL6* peptide treatment, and a stronger *GUS* signal was observed after the 5- μm peptide treatment than after the 1- μm treatment (Figure 4D). These results indicate that the *SIIDL6* signal may modulate *TAPG1*, *TAPG4*, and *CEL2* expression.

SIIDL6 and *PSK* regulate abscission via different pathways

A transcript analysis showed that both *SIPSKR1* (Solyc01g008140) and *SIPSKR2* (Solyc07g063000) were expressed during tomato pedicel abscission (Figure 5A). We also observed that *SIPSKR2* was significantly upregulated under low light stress (Supplemental Table S3; Figure 5B). Silencing *SIPSKR1* and *SIPSKR2* expression to inhibit the *PSK* signal significantly suppressed low light-induced abscission (Figure 5C). Furthermore, knockdown of *SIPSKR1* and *SIPSKR2* expression in the *CRslid6* lines further suppressed low light-induced abscission (Figure 5D), and similar results were observed after flower removal-induced abscission

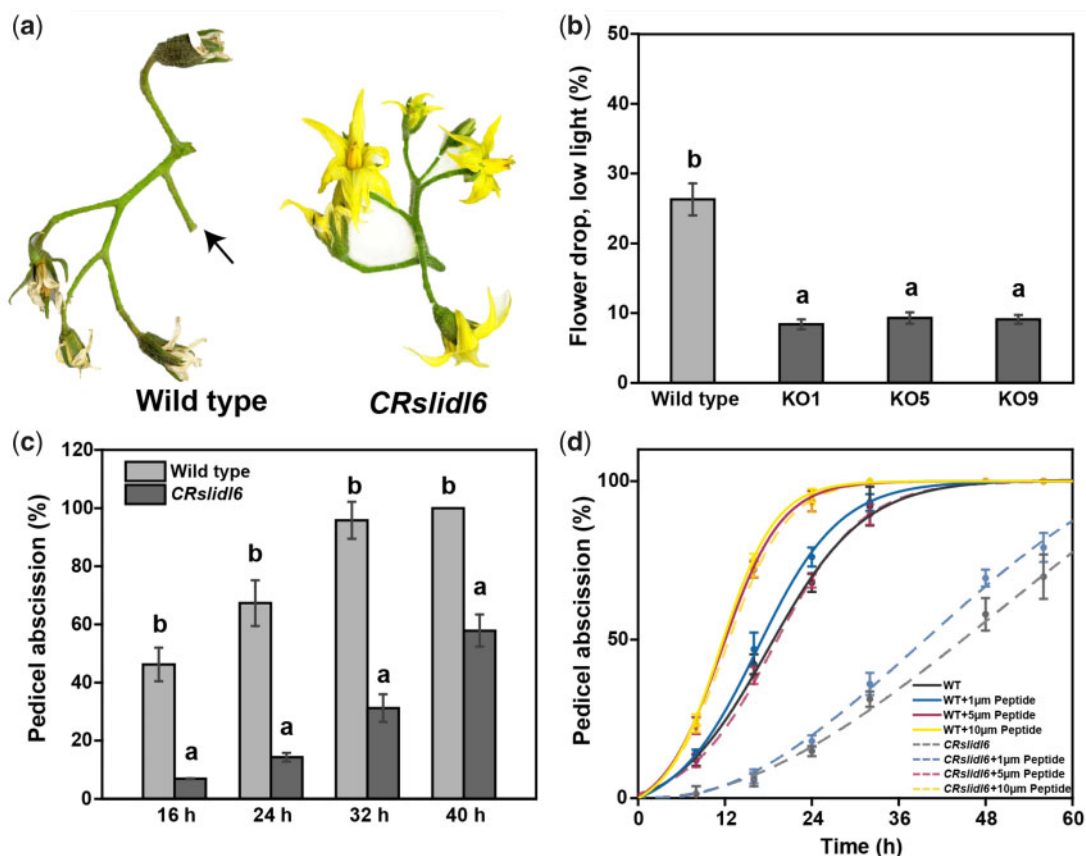


Figure 2 Effects of knocking out *SIIDL6* on the kinetics of flower pedicel abscission. A, Inflorescence phenotypes of *CRslid6* (knockout, KO) and the wild-type under low light stress. Wild-type shows premature abscission in the pedicel AZ (arrowhead). B, Low light-induced flower abscission in AZ (less than 0.5 mm) of *CRslid6* and wild-type lines after fruit set, and the percentage of flowers per inflorescence. The results are the means of three replicates \pm SD, with at least 18 samples per replicate. Different letters indicate significant differences (Student's *t* test, $P < 0.05$). C, The effect of flower removal on *CRslid6* and wild-type pedicel (about 4 cm) abscission. The results are the means of three replicates \pm SD, with at least 18 samples per replicate. Different letters indicate significant differences (Student's *t* test, $P < 0.05$). D, The effect of different peptide concentrations on *CRslid6* and wild-type pedicel abscission. The black, blue, purple, and yellow solid curves correspond to the abscission of wild-type lines after 0, 1, 5, and 10 μ m peptide treatment. The gray, blue, purple, and yellow dotted curves correspond to the abscission of *CRslid6* lines after 0, 1, 5, and 10 μ m peptide treatment. The results are the means of three replicates \pm SD, with at least 18 samples per replicate.

(Figure 5E). To further elucidate the relationship between *SIIDL6* and PSK signals during abscission, *SIIDL6*, PSK, and 5 or 12 mer control peptides were used to treat tomato pedicel explants. Co-treatment with *SIIDL6* and PSK led to significantly accelerated abscission compared with the single *SIIDL6* or PSK peptide treatments, whereas application of *SIIDL6* and the 5 aa control peptide, or PSK and the 12 aa control peptide, did not result in any differences compared with the single *SIIDL6* or PSK peptide treatments. These results indicate that *SIIDL6* and PSK act synergistically during abscission (Figure 5F).

SIIDL6 is directly regulated by *SIWRKY17*

To address the regulation of *SIIDL6* expression, a 1,284-bp promoter fragment upstream from its coding sequence was chosen. A search of the PlantCARE database (<http://bioinformatics.psb.ugent.be/webtools/plantcare/html/>) suggested that the promoter included two ABRE (ACGTG located at

–795 and –618) binding sites and one WRKY (GTCAGC located at –121) binding site (Figure 6A). A search of the low-light-treated tomato pedicel transcriptome data revealed a member of the WRKY II b subfamily, *WRKY17*, to be differentially expressed under low light conditions compared with wild-type light levels (Supplemental Table S3).

We investigated the binding of *SIWRKY17* to the *SIIDL6* promoter using a yeast one-hybrid assay and found that it bound and triggered the expression of the LacZ reporter gene (Figure 6B). To further confirm this interaction, we performed an electrophoretic mobility shift assay (EMSA) analysis using the full-length *SIWRKY17* protein and fragments of biotin-labeled *SIIDL6* promoter containing the W-box motif as well as a competitor probe containing three mutated nucleotides. We observed that *SIWRKY17* was bound to the W-box motif of *SIIDL6* (Figure 6C). Regulation of the *SIIDL6* promoter by *SIWRKY17* was investigated using a GUS transactivation assay in *Nicotiana benthamiana* leaves. An

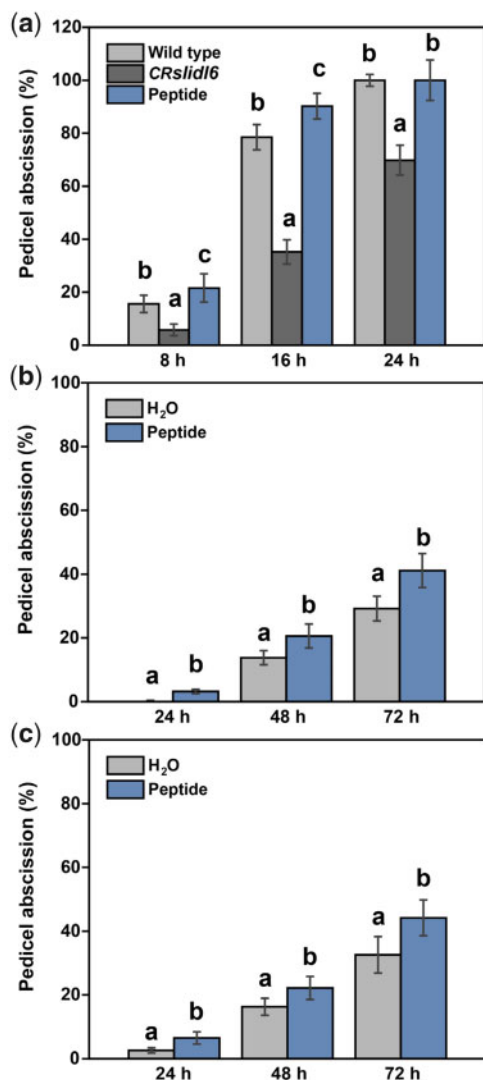


Figure 3 Kinetics of pedicel abscission is mediated by *SIIDL6*, which functions downstream from ethylene. A, Pedicel explant abscission assay for *SIIDL6* knockout (KO) and wild-type plants with or without 5- μ M peptide treatment and 10- μ L \cdot L $^{-1}$ ethylene. B, Pedicel explant abscission assay for 1-methylcyclopropene (1-MCP) pretreated wild-type pedicel explants with, or without, 5- μ M peptide. C, Pedicel explant abscission assay for *Never ripe* (*Nr*) mutant explants with, or without, 5- μ M peptide. Different letters indicate significant differences (Student's *t* test, $P < 0.05$). The results are the means of three replicates \pm SD, with at least 18 samples per replicate.

increase in the GUS activity was observed upon coexpression of 35S::WRKY17 with the wild-type *SIIDL6* promoter::reporter construct, but not for the mutated *SIIDL6* promoter (Figure 6D). The results indicated that *SIWRKY17* significantly promoted the activity of the *SIIDL6* promoter.

Knockdown of *SIWRKY17* delays pedicel abscission

We evaluated the expression of *SIWRKY17* in flower AZs after the low light treatments and observed elevated expressions in both cases, which we found to be blocked by treatment with 1-MCP (Figure 7A). Flower removal also induced *SIWRKY17* expression, which reached a peak at 24 h

after removal before decreasing (Figure 7B). Ethylene treatment induced *SIWRKY17* expression, while 1-MCP treatment suppressed its expression (Figure 7C).

Virus-induced gene silencing (VIGS) of *SIWRKY17* resulted in a significant downregulation of *SIWRKY17* but no downregulation of *SIWRKY6* (Solyc02g080890.2) or *SIWRKY16* (Solyc02g032950), while *SIIDL6* expression was lower in the pedicels of silenced tomato plants than in the control (Figure 7D; Supplemental Figure S5). Downregulation of *SIWRKY17* also decreased low light- and ethylene-induced *SIIDL6* expression (Supplemental Figure S6). The *SIWRKY17*-silenced flowers showed lower flower drop rates under low light stress and delayed pedicel abscission after flower removal (Figure 7, E and F).

Discussion

SIIDL6 is involved in low light-induced tomato flower abscission

Blossom drop is observed in many species following biotic or abiotic stress (Addicott, 1982; Taylor and Whitelaw, 2001; Tranbarger et al., 2017) and the IDA peptide is thought to play a conserved and central role in this process as well as in aspects of development (Shi et al., 2019). The expression of IDA and IDA-like genes is known to increase during abscission, and most have been shown to function in mediating organ abscission by controlling cell separation. In a previous study, ProAtIDA::GUS staining was not observed until the position 4 flower stage in *Arabidopsis*, but a strong signal in the floral AZ from positions 5 to 8 was seen, and an abscission-related expression pattern was observed. The overexpression of each of the genes *AtIDL1* to -5 was found to rescue the *ida* abscission defect (Butenko et al., 2003; Stenvik et al., 2008). Here, we found that *SIIDA*, *SIIDL6*, and *SIIDL7* were upregulated during pedicel abscission and that *SIIDL6* was induced by low light stress. We chose *SIIDL6* for further studies as it had the highest expression during pedicel abscission and low light stress; however, we cannot rule out the possibility that *SIIDA* and *SIIDL7* also contribute to tomato blossom drop (Supplemental Figure S7), which will be addressed in future studies.

The IDA peptide interacts with HAE and HSL2 to form a ligand-receptor module that triggers an abscission signal (Stenvik et al., 2008; Tang et al., 2017). A sequence alignment showed that *SIIDL6* does not have a PIPP sequence, but does contain a conserved core SGPS domain with a proline that may be post-translationally modified to form hydroxyproline. Notably, the presence of hydroxyproline residues was shown to increase the activity of CLAVATA3 (CLV3) and PLANT PEPTIDE CONTAINING SULFATED TYROSINE 1 (PSY1; Kondo et al., 2006). A positive effect of proline hydroxylation on peptide activity has also been reported for IDA (Butenko et al., 2014; Supplemental Figure S1). Furthermore, our results showed that *SIIDL6* plays a similar role to *AtIDA* in mediating floral abscission since its expression rescued the *ida2* mutant. In summary, our results show that the canonical IDA-HAE/HSL2 abscission signaling

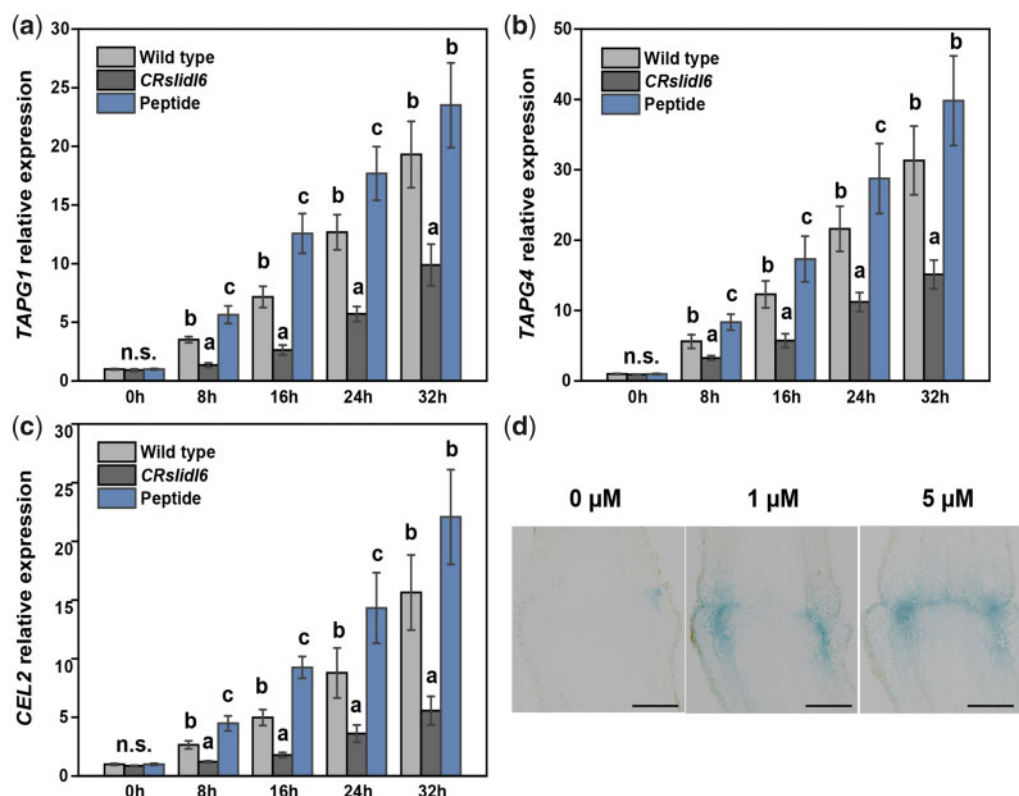


Figure 4 The expressions of genes encoding proteins involved in cell wall remodeling are affected by SIIDL6. RT-qPCR analysis of *TAPG1* (A), *TAPG4* (B), and *CEL2* (C) expressions in *CRslid6*, wild-type, and SIIDL6 peptide-treated plants. Different letters indicate significant differences (Student's *t* test, $P < 0.05$). The results are the means of three replicates \pm SD. D, β -glucosidase (GUS) staining revealed that the *TAPG4*::GUS signal is induced by different concentrations of SIIDL6 peptide 16 h after flower removal. Scale bar = 100 μ m.

found in *A. thaliana*, *Citrus*, and litchi is conserved in tomato (Aalen et al., 2013; Estornell et al., 2015; Ying et al., 2016).

SIIDL6 expression required ethylene but SIIDL6-induced abscission was partially ethylene independent

Ethylene is a well-known accelerator of flower abscission (Botton and Ruperti, 2019). In *A. thaliana*, the *ida* mutant showed a normal ethylene triple-response and senescence but not abscission following ethylene treatment, indicating that *ida* is not defective in ethylene signaling and that IDA signaling mediates organ abscission through an ethylene-independent pathway (Butenko et al., 2006). Recent studies have indicated that ethylene is required for mediating IDA expression and that wounding also induces IDA expression, although whether wounding directly induces IDA expression or acts via the induction of ethylene production is not clear (Meir et al., 2019). A previous study indicated that flower removal instantly depletes auxin levels and promotes AZ ethylene sensitivity (Meir et al., 2010). Our results showed that flower removal significantly induced *SIIDL6* expression after 4 h compared with flowers that remained attached to the explants, and that there was increased expression of AZ in the pedicel explants as abscission progressed. *SIIDL6* expression levels decreased after 1-MCP treatment and expression

in the *Nr* ethylene receptor mutant was depressed, indicating that *SIIDL6* expression is induced by ethylene. Low light-induced flower bud and flower abscission is influenced by ethylene action (Van Meeteren and De Proft, 1982; Wien et al., 1996); however, *SIIDL6* upregulation under low light stress is most likely ethylene dependent because 1-MCP suppressed low light stress-induced *SIIDL6* expression. We conclude that ethylene is partially dependent on *SIIDL6* when mediating abscission since abscission in *CRslid6* plants was significantly delayed, but not entirely blocked, following ethylene treatment. Furthermore, partially restored abscission in 1-MCP treated and *Nr* mutant explants after SIIDL6 peptide treatment, indicated that SIIDL6 could induce tomato pedicel abscission through a partially ethylene-independent pathway. As CLV3 and IDA peptides both carry hydroxyprolines, could be further modified by glycosylation, which resulting in enhancing peptide signaling (Ohyama et al., 2009). The FASCIATED INFLORESCENCE (*FIN*), which encodes a hydroxyproline O-arabinosyltransferase (HPAT). HPATs are enzymes that catalyze the transfer of L-arabinose to the hydroxyl group of hydroxyproline residues, and the mutant phenotype can be rescued by application tri-arabinosylated CLV3 peptides (Ogawa-Ohnishi et al., 2013; Shinohara and Matsubayashi, 2013). It is also reported the hydroxyprolination of IDA affects the affinity of the peptide for the HSL2 receptor (Butenko et al., 2014). We proposed

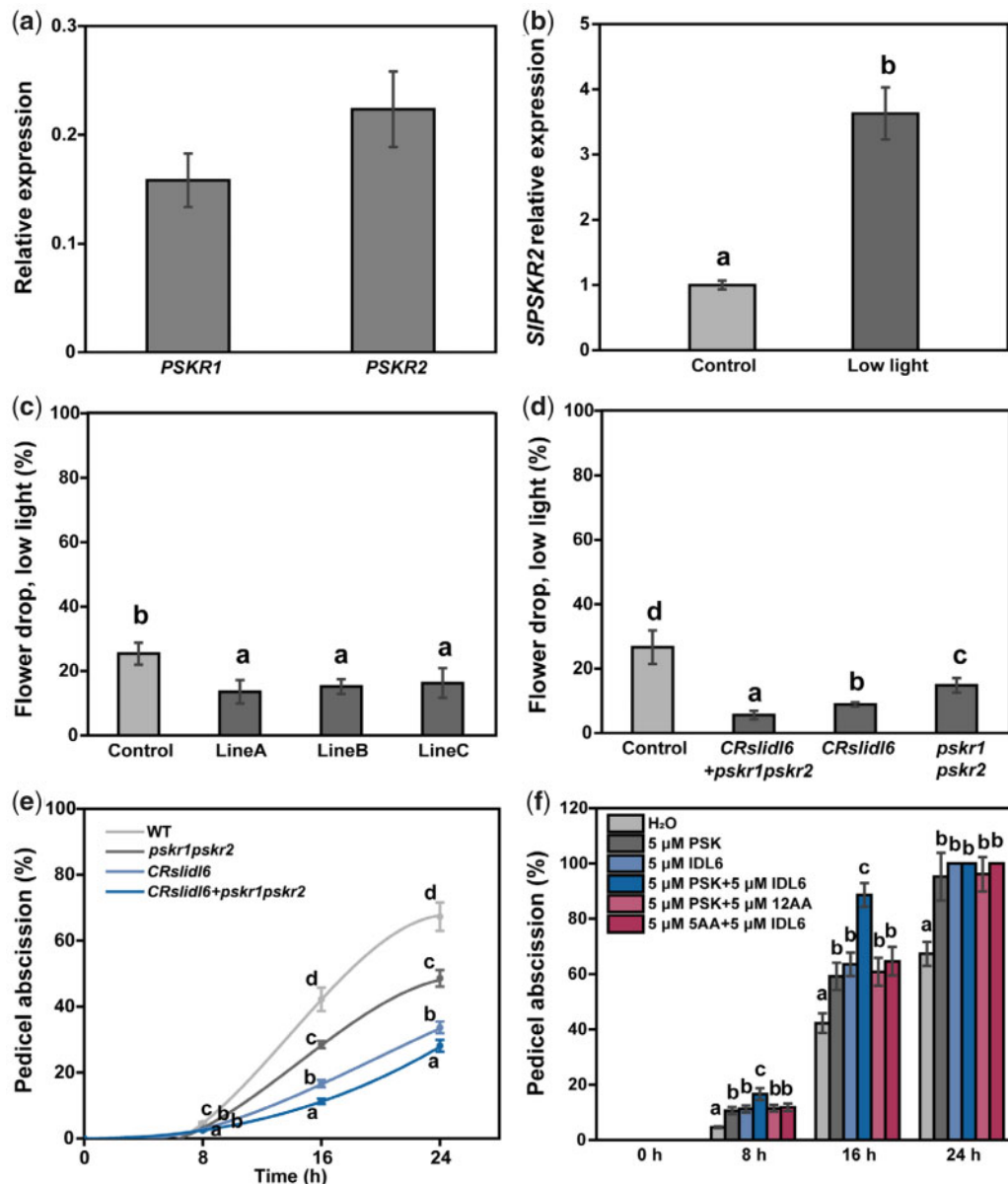


Figure 5 Effect of PSK and SIIDL6 peptide treatments on tomato flower pedicel abscission. A, *SIPSKR1* and *SIPSKR2* expression in the pedicel abscission zone. B, *SIPSKR2* expression in the pedicel abscission zone after 7 d of low light treatment compared with the control grown under normal light. The results are the means of three replicates \pm SD, with at least 20 samples per replicate (Student's *t* test, $P < 0.05$). C, Effect of *SIPSKR1* and *SIPSKR2* on low light-induced abscission. The results are the means of three replicates \pm SD, with at least 30 samples per replicate (Student's *t* test, $P < 0.05$). D, Effects of low light stress on pedicel abscission in pTRV0 control plants, *CRslid6* and *SIPSKR1*- and *SIPSKR2*-silenced pedicels of wild-type and *SIPSKR1*-, and *SIPSKR2*-silenced pedicels of *CRslid6* plants. The results are the means of three replicates \pm SD, with at least 30 samples per replicate (Student's *t* test, $P < 0.05$). E, The effect of flower removal on pedicel abscission in pTRV0 control plants, *CRslid6* and *SIPSKR1*-, and *SIPSKR2*-silenced pedicels of wild-type, and *SIPSKR1*- and *SIPSKR2*-silenced pedicels from *CRslid6* plants. The results are the means of three replicates \pm SD, with at least 30 samples per replicate (Student's *t* test, $P < 0.05$). F, Percentage abscised wild-type flower explants after exposure to 5- μ M PSK peptide, 5- μ M SIIDL6 peptide and both 5- μ M PSK peptide and 5- μ M SIIDL6 peptide, both 5- μ M PSK peptide and 5- μ M 12AA, and both 5- μ M 5AA peptide and 5- μ M SIIDL6 peptide compared with the control flowers. Different letters indicate significant differences between treatments (Student's *t* test, $P < 0.05$). The results are the means of three replicates \pm SD, with at least 30 samples per replicate.

that ethylene might regulate post-translational enzyme activities, which involved in hydroxyprolinated and glycosylated SIIDL6 peptide, to mediate abscission.

In *A. thaliana*, IDA is thought to act as a master regulator governing the final cell separation stages in the abscission process (Butenko et al., 2006; Stenvik et al., 2008; Aalen

et al., 2013). RNA-seq and microarray analyses have revealed that xyloglucan endotransglycosylase cell wall modification associated genes are significantly downregulated in both the *hae hsl2* and the *ida* mutant (Niederhuth et al., 2013; Reichardt et al., 2020). Similar to *A. thaliana*, knocking out *SIIDL6* suppressed *TAPG1*, *TAPG4*, and *CEL2* expression, but

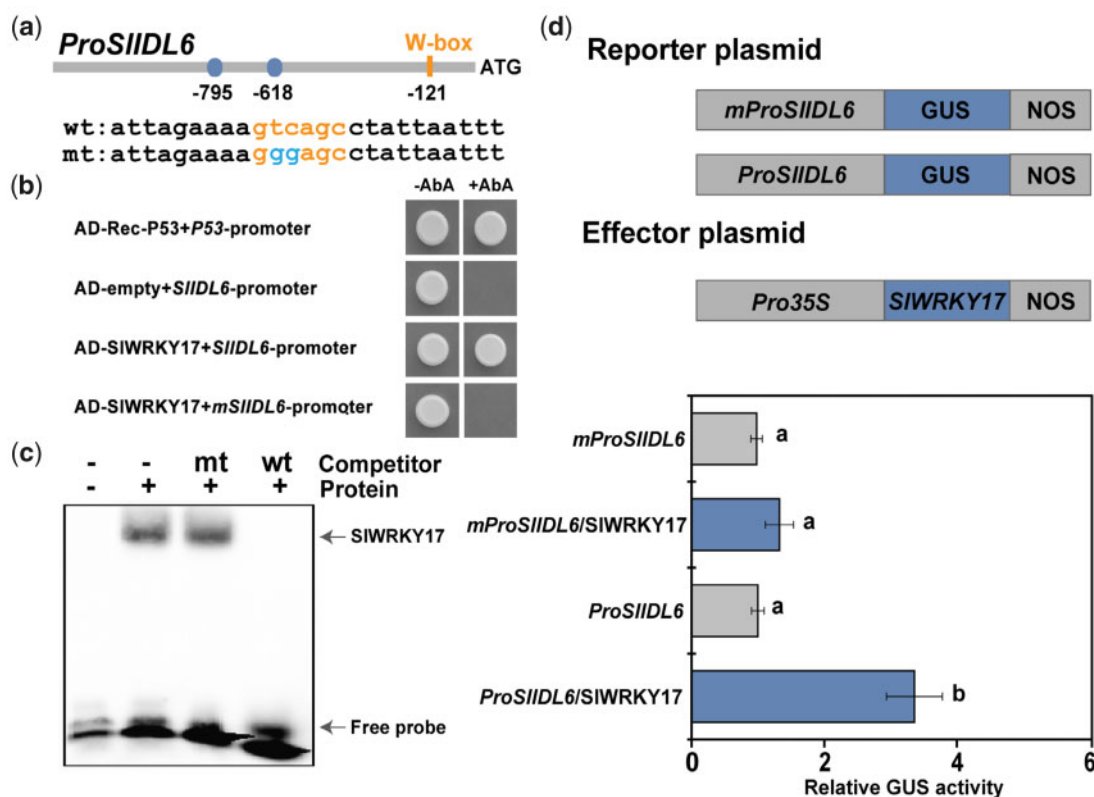


Figure 6 SIWRKY17 promotes SIIDL6 transcription. A, The SIIDL6 promoter fragment containing two ABRE (ACGTG located at -795 and -618) and one WRKY (GTCAGC located at -121) binding sites. wt, intact W-box element; mt, mutated W-box element. B, Yeast one-hybrid analysis revealing that SIWRKY17 binds to the SIIDL6 promoter fragment (Pro SIIDL6) containing the W-box motif (-121 bp). AbA (Aureobasidin A) was used as a screening marker. Rec-P53 and the P53-promoter were chosen as positive controls while the empty vector and the SIIDL6 promoter were used as negative controls. C, Electrophoretic mobility shift assay (EMSA) indicating that SIWRKY17 binds to the W-box motif of the SIIDL6 promoter. The biotin-labeled fragment of the SIIDL6 promoter containing the W-box motif (wt), a W-box motif with two mutated nucleotides (mt), and with a nonlabeled competitive probe (100-fold that of the biotin-labeled probe). His-tagged SIWRKY17 was used in the analysis. D, β -glucosidase (GUS) activity analysis showing that SIWRKY17 activates the SIIDL6 promoter. The SIWRKY17 overexpression vector combined with the reporter vector containing the wild-type SIIDL6 promoter, or a mutated promoter sequence, were co-infiltrated into wild *N. benthamiana* leaves to analyze the regulation of GUS activity. The transfection experiments were performed using three replicates. Values represent means \pm SD. Different letters indicate significant differences between treatments (Student's *t* test, $P < 0.05$).

applying SIIDL6 peptide had the opposite effect. It is highly acceptable that ethylene triggers IDA signaling, which in turn promotes the expression of cell wall hydrolases that break down polysaccharides in the cell wall and middle lamella, leading to abscission.

SIIDL6 and PSK signaling may regulate abscission in different ways

PSK signaling has been reported to be involved in drought-induced tomato flower abscission (Reichardt et al., 2020). PSK acts through an auxin- and ethylene-independent pathway to induce *TAPG4* expression and accelerate abscission. In tomato, the PSK peptide is perceived by both *PSKR1* and *PSKR2* (Zhang et al., 2018), and low light stress also induces the expression of *SIPSKR2*, suggesting a role in low light-induced abscission. Silencing *PSKR1* and *PSKR2* in tomato flowers resulted in delayed abscission under low light stress, indicating that PSK signaling is also involved in low light-

induced flower abscission. We investigated the relationship between SIIDL6 and PSK signaling because the SIIDL6 peptide treatment also induced *TAPG1* and *TAPG4* expression. PSK induced pedicel abscission, while the additional application of SIIDL6 has a synergistic effect on inducing abscission. Silencing *PSKR1* and *PSKR2* in the *CRslid6* lines further delayed abscission compared with the *CRslid6* lines. Given that SIIDL6 and PSK are both predominantly expressed in later abscission stages and are perceived by different receptors, we conclude that SIIDL6 and PSK signaling regulate abscission in different ways. Interestingly, AtIDA signaling has also been reported to be involved in drought-triggered leaf abscission, as the expression of the HAESA signaling receptor is induced in the leaf AZ under limited water conditions (Patharkar and Walker, 2016). In *N. benthamiana*, IDA-like genes also show induced expression under water stress (Ventimilla et al., 2020). During drought-induced tomato flower drop, *SIIDL1-5* is expressed at very low levels

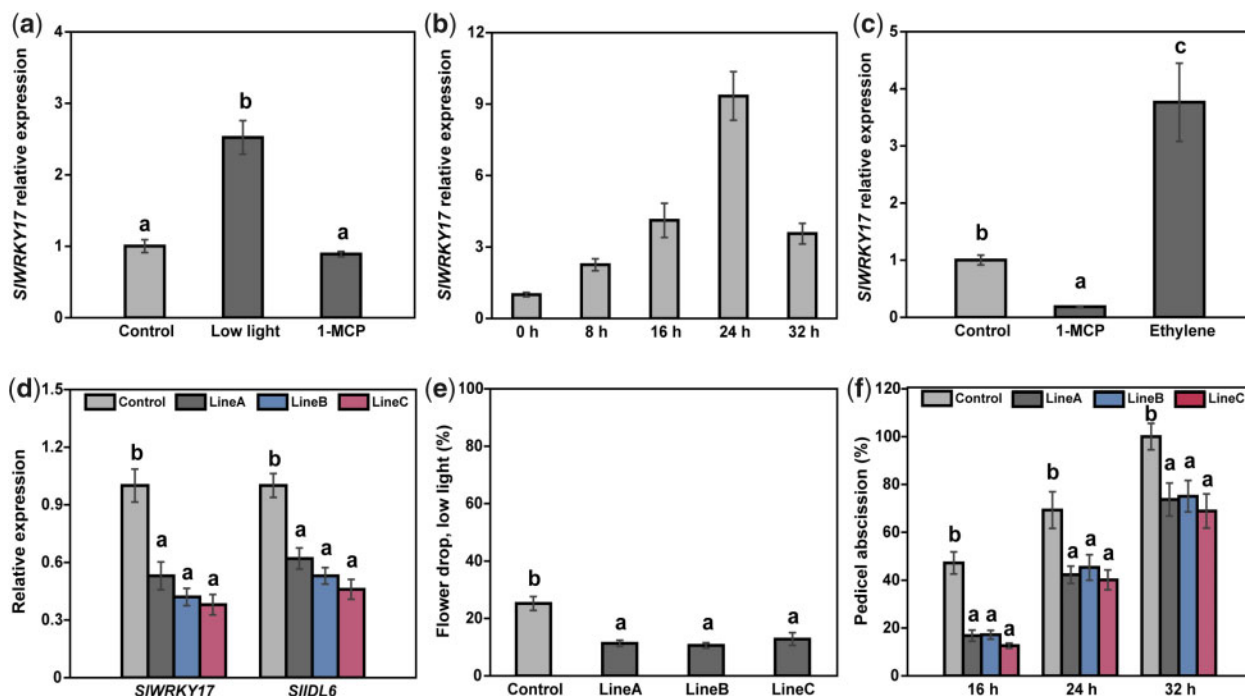


Figure 7 Silencing of *SIWRKY17* causes delayed flower abscission. A, *SIWRKY17* expression in the pedicel abscission zone (AZ) after 7 d of low light treatment compared with the control (normal light), and the effect of 1-methylcyclopropene (1-MCP) treatment on low light-induced *SIWRKY17* expression. B, Effect of flower removal on the *SIWRKY17* expression pattern in wild-type pedicel AZ. C, Effect of 10- $\mu\text{L}\cdot\text{L}^{-1}$ ethylene and 2- $\mu\text{L}\cdot\text{L}^{-1}$ 1-MCP on *SIWRKY17* expression in the pedicel AZ. D, Expression of *SIWRKY17* and *SIIDL6* in *SIWRKY17*-silenced and TRV control plants. E, Effect of low light stress on pedicel abscission in the TRV control and *SIWRKY17*-silenced plants. F, Effect of flower removal on *SIWRKY17*-silenced and wild-type pedicel abscission. Different letters indicate significant differences between treatments (Student's *t* test, *P* < 0.05). Results are the means of three replicates \pm SD.

(Reichardt et al., 2020). However, the role of *SIIDL6-11* and *SIIDA* in drought-induced abscission still needs to be elucidated. It may be that plants use multiple complex peptide-signaling systems to better control abscission and to coordinate the many associated cellular processes.

WRKY17 acts upstream of *SIIDL6* to trigger flower abscission

Based on their sequence, WRKY TFs have been grouped into six different classes (Zhang and Wang, 2005; Liang and Jiang, 2017). WRKY II b TFs have been shown to be involved in light signaling, stress responses, hormone signaling, leaf senescence, and abscission (Cheong et al., 2002; Meir et al., 2010; Koyama, 2014; Liang and Jiang, 2017). In this study, *SIWRKY17* was found to act as a positive regulator of *SIIDL6* by directly binding to the W-box elements of its promoter. *SIWRKY17* is also upregulated during low light-induced tomato flower abscission and ethylene treatment, and therefore probably plays a role in low light- and ethylene-induced abscission. The results of the EMSA, yeast one-hybrid assay, and GUS activity analyses were consistent with a regulatory *SIWRKY17*-*SIIDL6* module function in flower pedicel abscission. In tomato, *SIWRKY17* is upregulated by ethylene and is involved in ethylene signaling by mediating fruit ripening through its interaction with RIPENING INHIBITOR (SIRIN), ETHYLENE RESPONSE FACTOR 2b (SIERF2b), and

ETHYLENE RESPONSE FACTOR 7 (SIERF7), and silencing *SIWRKY17* caused reduced red fruit coloration (Wang et al., 2017). Furthermore, *SIWRKY17* knockdown suppressed low light stress- and ethylene-induced *SIIDL6* expression and delayed abscission during low light stress and ethylene treatment, which is consistent with the phenotypes of the *CRslid6* lines. We propose that low-light stress enhances ethylene signaling to trigger the *SIWRKY17*-*SIIDL6* regulatory module, which induces flower pedicel abscission (Figure 8).

Materials and methods

Plant materials

Transgenic tomato plants were generated using the wild-type *Solanum lycopersicum* cv Ailsa Craig (AC), and *TAPG4::GUS* seeds were provided by Prof. Mark L. Tucker (Hong et al., 2000). The wild-type, transgenic, and tomato ethylene response mutant, Nr seeds were washed in 50% (v/v) bleach for 3 min, rinsed, and placed onto paper soaked in water in culture dishes for 3 d at 28°C. Then the seedlings were grown in soil for seven weeks in a greenhouse under standard greenhouse conditions (25°C under 16 h of light, followed by 8 h of dark at 15°C).

Tomato plants were subjected to low light conditions in the greenhouse by covering them with shading nets. The photosynthetically active radiation levels under normal and shade conditions were 600 and 60 μmol , respectively. For

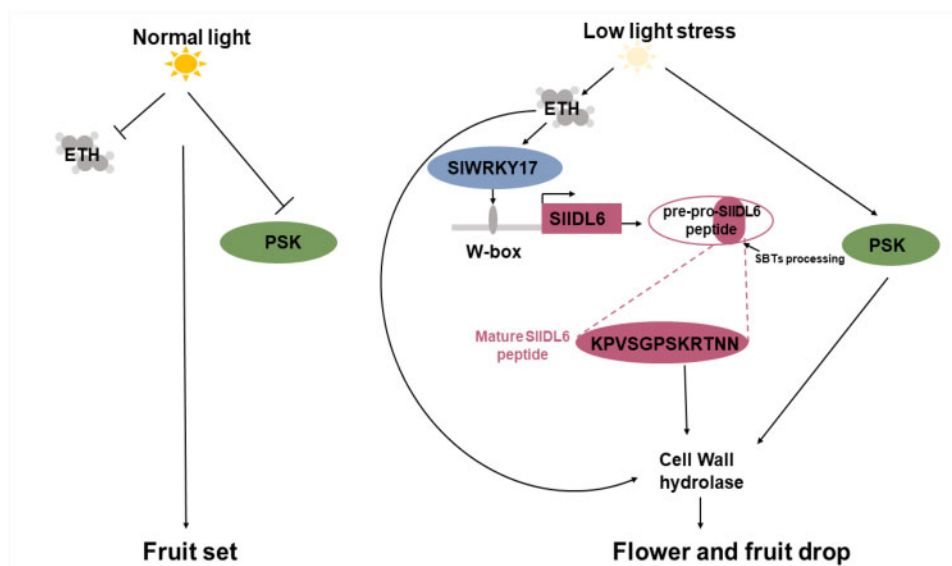


Figure 8 Model of SIIDL6 signaling during low light-induced tomato pedicel abscission. Normal light guarantees fruit set by inhibiting ethylene signaling and PSK expression. Low light stress enhances ethylene signaling, which activates *SIWRKY17* to induce *SIIDL6* expression in the AZ. The mature SIIDL6 peptide is secreted and acts as a signal to promote expression of cell wall hydrolases, leading to flower drop. ETH, ethylene; PSK, phytosulfokine; SBT, subtilisin-like proteinase; W-box, WRKY binding site.

the ethylene treatments, tomato flower explants were incubated in 0.9% agar in a $40 \times 25 \times 20$ cm³ glass container, which was injected with ethylene to a final concentration of $20 \mu\text{L L}^{-1}$. In the 1-MCP pretreatments, plants were incubated with 1-MCP in a sealed 200 L chamber at a final concentration of $0.4 \mu\text{L L}^{-1}$ at 22°C for 8 h. The cumulative abscission rates of the tomato pedicels were determined as previously described (Wang et al., 2005).

Plasmid construction and plant transformation

The *SIIDL6* (Solyc06g050140) was overexpressed by amplifying the full-length *SIIDL6* cDNA using gene-specific primers. The full-length fragment was cloned into the pENTR/D-TOPO vector (Invitrogen) and then cloned into the binary pB7YWG2 vector using BP and LR clonase (Invitrogen), creating the 35S::SIIDL6 plasmid. To make deletions in the *SIIDL6* coding sequence and produce knockout lines, a construct (*CRslid6*) was designed that contained two single-guide RNAs (sgRNAs) and the Cas9 endonuclease gene. Two single-guide RNAs (sgRNA1 and sgRNA2; Supplemental Figure S2), were designed to target the *SIIDL6* gene, using the CRISPR-P tool (<http://cbi.hzau.edu.cn/crispr>). The U6-26-SIIDL6-gRNA cassettes were cloned into the CRISPR/Cas9 binary vector, pCBC-DT1T2_tomatoU6, to generate pCBC-DT1T2_tomatoU6-SIIDL6.

The *CRslid6* and 35S::SIIDL6 vectors were introduced into *Agrobacterium tumefaciens* LBA4404 cells by electroporation and then transformed into wild-type tomato by leaf disc cocultivation (Wang et al., 2018), or into the *A. thaliana* *ida2* mutant using the floral dip technique. Positive *CRslid6* lines were identified by sequencing PCR products amplified from the targeted region. Cas9-free T2 plants carrying mutations were screened and identified. Positive *A. thaliana*

35S::SIIDL6 lines were obtained by spraying them with a 250-mg L^{-1} Basta solution two to three times and confirmed by PCR. All primers used are listed in Supplemental Table S1.

Virus-induced gene silencing

A 325 bp *SIWRKY17* (Solyc07g051840) specific fragment (from 809 to 1,133 bp) was amplified by PCR and purified using the TaKaRa PCR fragment recovery kit before being cloned into the pMD-18T T-vector and verified by sequencing. The *EcoRI* restriction enzyme site was then used to insert the fragment into the tobacco rattle virus pTRV2 vector. The pTRV1 and pTRV2-SIWRKY17 vectors were individually transformed into *A. tumefaciens* strain GV3101. A mix of the two *A. tumefaciens* cultures were used to infiltrate the tomato leaves.

Phylogenetic analysis

A BLASTp search was performed with the conserved amino acid sequences from AtIDA and AtIDL1-8 against the tomato genomic database (<https://www.solgenomics.net/>). Phylogenetic analysis of SIIDA amino acid sequences (SIIDA [Solyc05g010000.1]; SIIDL1 [CP023762.1]; SIIDL2 [CP023760.1]; SIIDL3 [Solyc07g044890.1]; SIIDL4 [Solyc05g007040.1]; SIIDL6 [Solyc06g050140.1]; and SIIDL7 [Solyc09g005780.1]) with AtIDAs amino acid sequences was performed using the MEGA5 software with the neighbor-joining method (Tamura et al., 2011).

In situ hybridization

The different abscission stages of the wild-type pedicel explants were sampled and immediately fixed in phosphate-buffered saline solution containing 4% (w/v) paraformaldehyde. After submersion in an ethanol gradient for

dehydration, the samples were embedded in Paraplast Plus (Leica) and sectioned into 10 μm slices using a microtome (Leica). In situ hybridization analysis of *SIIDL6* was performed as previously described (Wang et al., 2018). The DIG Oligonucleotide 3'-End Labeling Kit (Roche) was used to generate antisense and sense RNA probes. Probes that contained the unique *SIIDL6* region were 76 bp long (from 174 to 249 bp).

RNA extraction and RT-qPCR analysis

TRIzol reagent (Invitrogen) was used to extract RNA from different tomato pedicel AZ samples. The RNA was treated with recombinant DNase I (TaKaRa) according to the manufacturer's instructions. After removing the DNA, 2- μg RNA (20- μL reactions containing 4- μL 5X PrimeScript RT Master Mix) was used to synthesize first-strand cDNA with an oligo (dT) primer and reverse transcriptase (TaKaRa). RT-qPCR was performed in 20- μL reactions containing 1 μL of cDNA, 200 nM of each primer and SuperReal PreMix Plus (Tiangen) on an Applied Biosystems 7500 instrument (Applied Biosystems). The mRNA and *SIIDL6* were detected using amplification reactions and conditions that had been set according to the manufacturer's protocols for SYBR SuperReal PreMix plus. Actin (NCBI: NM_001330119.1) was used as a control for mRNA. The program of RT-qPCR was 95°C 30 s, 95°C 5 s (40 cycles), 60°C 34 s (40 cycles). The RT-qPCR experiments were performed on three biological samples with three technical replicates. All primers used are listed in Supplemental Table S1.

Peptide treatment

Wild-type tomato flower pedicel explants were obtained at anthesis rinsed briefly in 70% ethanol and distilled water, and then incubated on 0.9% agar plates containing 0, 1, 5, or 10 μM concentrations of the *SIIDL6* peptide or 0, 1, and 5 μM of the sulfated PSK peptide, or both, with or without 5AA(TNGTK) and 12AA(PFLGVYYHKNNK) control peptide substitutes. Chromatography water was used as a control. The plates were placed under a glass cover in a growth chamber at 25°C for 8 h in the dark and under 16 h of light. Data were collected and analyzed from three independent treatments. The peptides were synthesized by the Shengggong Company (China), the *SIIDL6* had a purity of > 90%, and the PSK had a purity of >80%.

pBS assay

The force needed to remove the petals from 2, 4, and 6 positions on the inflorescence was quantified using pBS, as previously described (Stenvik et al., 2008). Data were collected and analyzed from at least 15 plants at each position.

Electrophoretic mobility shift assay

The conserved domains of *SIWRKY17* (631 amino acids, aa) were cloned and inserted into the pEASY-E1 vector (Transgene) containing a His tag sequence and then transformed into *Escherichia coli* BL21 cells (Transgene). The transformed cells were cultured in LB medium containing 50

$\mu\text{g mL}^{-1}$ of ampicillin at 37°C for $\text{OD}_{600} = 0.5$, and a final concentration of 0.2-mM isopropyl β -D-1-thiogalactopyranoside (IPTG, TaKaRa) was used to induce protein expression after incubation for 16 h at 23°C. Purification of the recombinant fusion proteins was performed. The 3' biotin end-labeled DNA probes were prepared by the Shengggong Company (China). A Light Shift Chemi Luminescent EMSA Kit (Thermo Scientific) was used to perform the EMSA assay. Nonmutated or mutated probes (20 fmol final concentration) were added to a solution of 2% glycerol, 50-mM KCl, and 10-mM EDTA buffer, with or without *SIWRKY17* protein, and incubated for 10 min at 23°C.

LC-MS/MS analysis

After removal of the flower, 0, 16, and 32 h pedicel AZs were collected and about 600 AZs were pooled respectively. The peptides were extracted by homogenization in 10 mL of phenol saturated with 50-mM Tris-HCl (pH 8.0) and precipitated with acetone. Size exclusion chromatography and reversed-phase chromatography were used to concentrate the samples. The Huada Protein Research Center (Beijing, China) performed the LC-MS/MS analysis, and Proteome Discoverer 2.1 software (Thermo Fisher Scientific) was used to analyze the data by searching against *SIIDL6* peptide databases, which contained 1,140 sequences, ranging in length from 10 to 22 amino acids to ensure that all possible peptides were identified. The scan selected two to eight charged state ions in positive mode with a minimum and maximum precursor mass of 350 and 5,000 Da, respectively. The threshold values of the precursor and fragment mass high-energy collision dissociation used for the search were 10 ppm and 0.08 Da, respectively.

Yeast one-hybrid assay

The full-length *SIWRKY17* sequence was ligated into the pGADT7 vector and the *SIIDL6* promoter fragment was cloned into the pAbAi vector. The MatchmakerTM Gold Yeast One-Hybrid Library Screening System kit was used to conduct a yeast one-hybrid assay, according to the manufacturer's instructions.

β -glucuronidase activity assay

The effector construct was obtained by cloning the coding sequence (CDS) of *SIWRKY17* (1–957 bp) into the binary pRI101 vector using the *HindIII* and *EcoRI* enzymes. To generate reporter constructs, the *SIIDL6* promoter sequence (1,284 bp) was cloned into the binary pBI101 vector, which contains the GUS reporter gene. The reporter and effector vectors were transformed into *A. tumefaciens* EHA105 and co-infiltrated into *N. benthamiana* leaves. Infected leaves were incubated in a growth chamber for 3 d. Then the proteins were extracted. A BCA Protein Assay Kit (Beyontian) was used to determine protein concentration, and GUS activity was determined from three biological replicates and leaves obtained at time zero were used as a control.

Accession numbers

Sequence data from this article can be found in the Genome Database for Tomato (<https://www.solgenomics.net/>) or GenBank/EMBL libraries under accession numbers SIIDA (Solyc05g010000.1); SIIDL1 (CP023762.1); SIIDL2 (CP023760.1); SIIDL3 (Solyc07g044890.1); SIIDL4 (Solyc05g007040.1); SIIDL6 (Solyc06g050140.1); and SIIDL7 (Solyc09g005780.1).

Supplemental data

The following materials are available in the online version of this article.

Supplemental Table S1. Primer sequences.

Supplemental Table S2. SIIDA/SIIDLs DNA and protein sequences.

Supplemental Table S3. IDL6, PSKR2 and WRKY17 expression in tomato flower pedicels AZ under low light.

Supplemental Table S4. Determination of mature peptides.

Supplemental Table S5. Potential off-target sites in CRISPR mutants.

Supplemental Figure S1. IDA/IDL protein sequence analysis using *Arabidopsis thaliana* and tomato sequences.

Supplemental Figure S2. Sequences of the eight *CRslid6* lines.

Supplemental Figure S3. Overexpression of *SIIDL6* in floral AZ of the *Arabidopsis thaliana ida2* mutant.

Supplemental Figure S4. Petal break strength in *ida2* and *ida2 35S::SIIDL6* plants.

Supplemental Figure S5. Expressions of the *SIWRKY6* and *SIWRKY16* genes in *SIWRKY17*-silenced and TRV control plants.

Supplemental Figure S6. Silencing of *SIWRKY17* causes decreased *SIIDL6* expression.

Supplemental Figure S7. RT-qPCR analysis of *SIIDLs* expression.

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Conflicts of interest: The authors declare no conflicts of interest.

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