

The Pentatricopeptide Repeat Protein SOT5/EMB2279 Is Required for Plastid *rpl2* and *trnK* Intron Splicing^{1[OPEN]}

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Chloroplast biogenesis and development are highly complex processes requiring interaction between plastid and nuclear genomic products. Using a high-throughput screen for chloroplast biogenesis suppressors in *Arabidopsis* (*Arabidopsis thaliana*), we identified a suppressor of *thf1* (*sot5*) that displays virescent and serrated leaves. Further characterization revealed that *sot5* mutants are defective in leaf adaxial and abaxial polarity and act as enhancers of *asymmetric leaves2*. Map-based cloning identified SOT5 as a gene previously named EMB2279 that encodes a plastid-targeted pentatricopeptide repeat (PPR) protein with 11 PPR motifs. A G-to-A mutation in *sot5* leads to a significant decrease in splicing efficiency, generating two additional mRNA variants. As reported previously, the *sot5* null mutation is embryo lethal. SOT5 is predicted to bind to specific RNA sequences found in plastid *rpl2* and *trnK* genes, and we found decreased splicing efficiency of the *rpl2* and *trnK* genes in *sot5* mutants. Together, our results reveal that the PPR protein SOT5/EMB2279 is required for intron splicing of plastid *rpl2* and *trnK*, providing insights into the role of plastid translation in the coupled development between chloroplasts and leaves.

Chloroplast development, which is an essential event for the switching of plant growth from heterotrophy to autotrophy, relies on the coordinated expression of nuclear and plastid genes. Although significant progress has been made in the past few years toward understanding the molecular mechanisms underlying chloroplast development, it remains elusive how chloroplast gene expression is finely regulated in response to intracellular and environmental cues (Pogson et al., 2015; Chan et al., 2016). Since chloroplasts originated from endosymbiotic cyanobacteria and possess prokaryote-type circular DNA, the chloroplast remains the same machinery that carries out gene expression as prokaryotes (Jarvis and López-Juez, 2013). However, some novel events, such as RNA editing, intron splicing, and RNA cleavage and trimming (Stern et al.,

2010), have evolved, making gene expression in chloroplasts distinct from that in prokaryotes.

In *Arabidopsis* (*Arabidopsis thaliana*) chloroplasts, 20 introns are distributed across 17 genes (de Longevialle et al., 2010); 19 of these introns belong to the group II family of introns, and only one, located in the *trnL* gene, belongs to group I. Original group I and II introns are large ribozymes that cleave phosphodiester bonds through two consecutive trans-esterification steps in a sequence-specific manner (Lehmann and Schmidt, 2003). The main differences between these two types of introns lie in their secondary structures and chemical reaction mechanisms for the first splicing step: group I introns are composed of 10 domains (P1–P10) that play a role in RNA folding for efficient splicing and ligation; group II introns have six domains (I–VI), each with distinct functions in intron self-splicing and/or retromobility (Lehmann and Schmidt, 2003; Dai et al., 2008; Pyle, 2016). The largest domain, domain I, functions as a structural scaffold and recognizes the exon positions for catalysis. Domains II and III enhance the catalysis of splicing. Domain IV, the most variable domain in different introns, contains the intron-encoded protein (IEP) sequence, which constitutes four different subdomains (reverse transcriptase, maturase, DNA binding, and endonuclease) that are required for RNA folding during splicing and intron retromobility after splicing. Chloroplast introns have no IEP sequence except for the *trnK* intron, which contains a limited coding sequence only for functional maturase (MatK; Zoschke et al., 2010) and the break point of the trans-splicing *rps12-1* intron (Glanz and Kück, 2009). The highly conserved domain V contains the catalytic core that is essential for ribozyme activity. Domain VI

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contains the branch-point site adenosine required for the splicing reaction. Group II introns are further classified into several subclasses (IIA–IIF) that display distinct structural and functional differences (Pyle, 2016). Among the 19 group II introns in Arabidopsis chloroplasts, eight belong to group IIA and 11 belong to group IIB (de Longevialle et al., 2010). However, both group I and group II introns in plant organelles are degenerated and require additional trans-factors for efficient splicing in vivo (Hahn et al., 1998; de Longevialle et al., 2010). These trans-factors, including chloroplast IEP (MatK) and nucleus-encoded RNA-binding factors, are assembled with cis-elements (intron RNAs) into active ribonucleoprotein complexes for accurate and efficient self-splicing (Stern et al., 2010; Zoschke et al., 2010). The chloroplast *matK* gene was reported to be required for the splicing of seven group IIA introns, *trnA*, *trnI*, *trnV*, *atpF*, *rpl2*, *rps12-2*, and *trnK* (Zoschke et al., 2010).

In addition to the trans-factor encoded by *matK*, a number of nucleus-encoded proteins involved in chloroplast splicing events have been identified, particularly from maize (*Zea mays*), through genetic and biochemical approaches. Some of the trans-factors are required for splicing multiple introns, while others are required for specific introns (Stern et al., 2010). Chloroplast RNA Splicing1 (CRS1) and CRS2 (Jenkins et al., 1997) were the first reported trans-factors in maize. Whereas CRS1 contains three repeated chloroplast RNA splicing and ribosome maturation (CRM) domains and is essential solely for the splicing of the chloroplast *atpF* group II intron (Till et al., 2001; Keren et al., 2008), CRS2 is a homolog of bacterial peptidyl-tRNA hydrolase and required for the splicing of nine group II introns (Jenkins et al., 1997; Ostheimer et al., 2003). CRS2 forms a complex with CRS2-Associated Factor1 (CAF1) and/or CAF2, both of which are CRM proteins with two CRM domains, to facilitate the splicing of the CRS2-dependent introns (Ostheimer et al., 2003). The splicing specificity of CRS2/CAF-dependent introns is determined by CRM Family Member2 (CFM2) and CFM3 (Asakura et al., 2008). Two general splicing factors, an RNase III domain protein (RNC1) and WTF1 (WHAT IS THIS FACTOR1), were identified from CAF1- and/or CAF2-immunoprecipitated ribonucleoprotein particles by mass spectrometry (Watkins et al., 2007). RNC1, which contains two RNase III domains without RNA cleavage activity, and WTF1, which has the plant organelle RNA recognition domain (PORR), form heterodimers that splice most of chloroplast group II introns in a CAF1/CAF2-dependent or -independent manner (Kroeger et al., 2009). It remains unclear whether a core spliceosome, as in the nucleus, is present for the in vivo splicing of all group II introns in chloroplasts.

In addition to these general splicing factors, another group of proteins involved in chloroplast intron splicing events are pentatricopeptide repeat (PPR) proteins, which bind to their single-stranded RNA targets in a modular and base-specific fashion (Yin et al., 2013;

Barkan and Small, 2014). For example, PPR4 and PPR5 are responsible for the splicing of the *rps12* and *trnG-UCC* group II introns in maize, respectively (Schmitz-Linneweber et al., 2006; Beick et al., 2008); in Arabidopsis, ORGANELLE TRANSCRIPT PROCESSING70 (OTP70) is required for the splicing of the *rpoC1* intron (Chateigner-Boutin et al., 2011), OTP51 for *ycf3* intron 2 and several other group IIA introns as well (de Longevialle et al., 2008), THYLAKOID ASSEMBLY8 for *ycf3* intron 2 and *trnA* (Khrouchtchova et al., 2012), EMBRYO DEFECTIVE2654 (EMB2654) for trans-splicing of *rps12* intron 1 (Aryamanesh et al., 2017); in rice (*Oryza sativa*), WHITE STRIPE LEAF (WSL) is required for *rpl2* intron splicing (Tan et al., 2014) and WSL4 for four group II introns, *atpF*, *rpl2*, *rps12-2*, and *ndhA* (Wang et al., 2017). Generally, PPR proteins are characterized by a one-repeat/one-nucleotide recognition mechanism (Prikryl et al., 2011; Yin et al., 2013). Thus, the binding specificity and affinity of a PPR protein to RNA targets are associated with the number of PPR repeats. It has been known that over 450 PPR proteins are present in Arabidopsis and divided into two subfamilies, designated P and PLS, according to the characteristics of the PPR motifs (Lurin et al., 2004). Most PPRs are involved in organellar gene expression, including RNA transcription and stability, RNA editing, RNA maturation, RNA translation, and RNA splicing (Schmitz-Linneweber and Small, 2008). However, the roles of most PPR proteins in RNA processing still remain unknown. It is thought that, during the group II intron splicing process, the intron-specific PPRs act in an initial sequence-specific binding step that alters RNA conformation ready for the recruitment of the relevant general splicing complex, such as the WTF/RNC1 complex (de Longevialle et al., 2010). This hypothesis still remains to be validated by experimental data.

Variation mutants are valuable for the study of chloroplast biogenesis mechanisms. The study of suppressors of variegation mutants, such as *var2* (*variegation2*) and *thf1* (*thylakoid formation1*), has provided lots of insights into the mechanisms of chloroplast development (Yu et al., 2008; Wu et al., 2013, 2016; Hu et al., 2015; Ma et al., 2015). Here, we report another suppressor of *thf1*, namely *sot5* (*suppressor of thf1 5*), whose gene encodes one of the Arabidopsis PPR proteins, EMB2279. We found that EMB2279/SOT5 is required specifically for plastid *rpl2* and *trnK* intron splicing. The *sot5* defects in plastid translation and leaf morphogenesis may indicate a role of plastid gene expression in regulating leaf development.

RESULTS

Identification and Characterization of the *sot5* Mutant

To investigate the molecular mechanisms by which *THF1* mutations lead to defects in chloroplast development, we mutagenized *thf1* seeds with ethyl

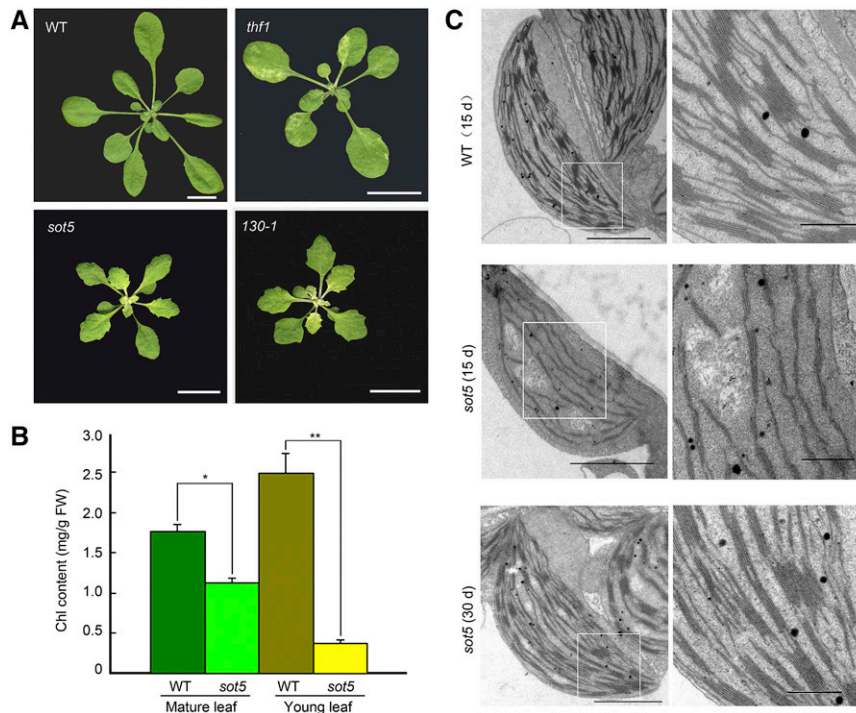


Figure 1. Identification and characterization of the *sot5* mutant. A, Phenotypes of 30-d-old wild-type (WT), *thf1*, *sot5*, and *130-1* plants. Bars = 1 cm. B, Chlorophyll contents of mature and young leaves from 45-d-old wild-type and *sot5* plants. The data represent means \pm SD of three biological replicates. Asterisks indicate significant differences between the wild type and *sot5* (Student's *t* test, *, $P < 0.05$ and **, $P < 0.01$). FW, Fresh weight. C, Ultrastructure of chloroplasts in the first wild-type and *sot5* true leaves sampled from 15- or 30-d-old plants. The images at right are closeups of the boxes in the left images. Bars = 2 μ m at left and 0.5 μ m at right.

methanesulfonate and screened for suppressors of *thf1* leaf variegation, as reported previously (Wu et al., 2013, 2016; Hu et al., 2015; Ma et al., 2015). The *sot5* single mutant was isolated from the F2 generation of the cross between a suppressor line, *130-1*, and Columbia-0 (Col-0). Both *sot5* and *130-1* displayed the same phenotype, such as virescent and serrated leaves (Fig. 1A). *sot5* cotyledons were yellowish and contained about 74% of the chlorophyll (Chl) content of wild-type cotyledons (Supplemental Fig. S1, A and B). By contrast, young true leaves of *sot5* were initially pale yellow but gradually became light green (Fig. 1A; Supplemental Fig. S1C). Total Chl analysis showed that *sot5* had about 15% and 64% of the Chl content of the wild type in young and mature leaves, respectively (Fig. 1B). Transmission electron microscopy analysis of chloroplast ultrastructure demonstrated that chloroplasts in *sot5* leaves were initially smaller and had fewer thylakoid membranes than those in wild-type leaves (Fig. 1C), but this difference was not apparent in mature leaves (Fig. 1C). Taken together, these results indicate that *SOT5* is required for chloroplast development.

We also analyzed the photosynthetic capacity of *sot5*. The F_v/F_m value, which represents the maximum quantum yield of PSII photochemistry, was significantly lower in young leaves of *sot5* than those of the wild type but was comparable in the mature leaves between

sot5 and the wild type (Fig. 2, A and B). NPQ, which is an important photoprotective mechanism to dissipate excessive light energy absorbed by antennae as heat, increased more rapidly in *sot5* than in the wild type in both mature and young leaves, whereas there was no obvious difference in NPQ relaxation in the dark between *sot5* and the wild type (Fig. 2B), and young *sot5* leaves maintained a higher level of NPQ than wild-type leaves in light (Fig. 2B). Immunoblot analysis showed that all photosynthetic proteins examined, such as D1, Lhca1, Lhcb6, PsdA, and RbcL, accumulated to apparently significantly lower levels in *sot5* young leaves (Fig. 2C) compared with the wild type, but the difference of these proteins between the wild type and *sot5* disappeared in mature leaves. Further analysis of photosynthetic complexes in *sot5* leaves using blue-native (BN) PAGE assays showed that *sot5* contained the same pattern of photosynthetic complexes as the wild type (Fig. 2D). Together, these results indicate that loss-of-function *SOT5* reduces photosynthetic efficiency and capacity.

Leaf Morphology and Cell Proliferation Were Altered in *sot5*

In addition to photosynthesis-related defects, *sot5* mutants displayed leaf morphology defects. The first

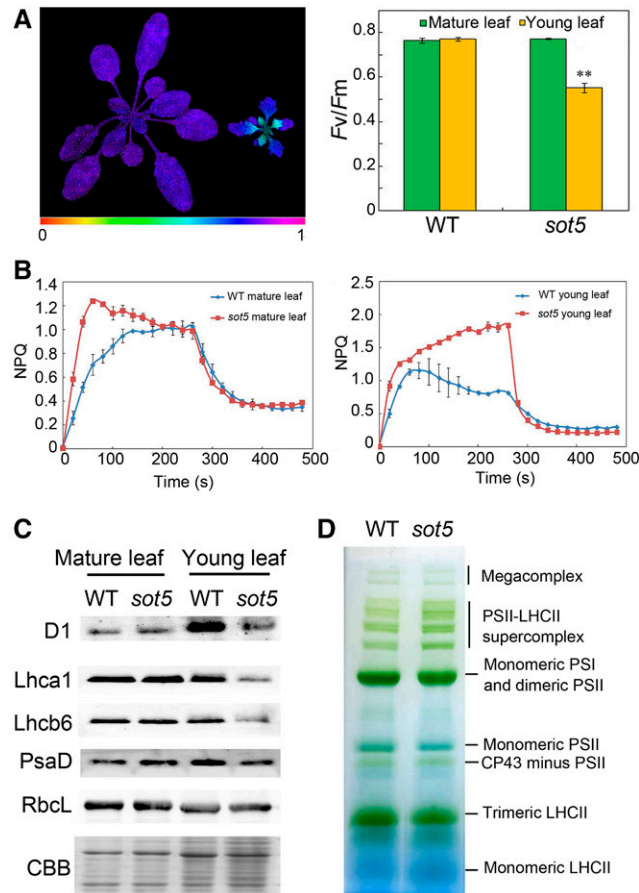


Figure 2. Photosynthetic characterization of *sot5* mutants. **A**, Analysis of F_v/F_m of 30-d-old wild-type (WT) and *sot5* plants. The graph at right shows quantified F_v/F_m of the mature and young leaves shown at left (Chl fluorescence image). The data represent means \pm SD ($n = 6$). Asterisks indicate a significant difference between the wild type and *sot5* (Student's *t* test, **, $P < 0.05$). **B**, Nonphotochemical quenching (NPQ) measurement of mature and young leaves from 30-d-old wild-type and *sot5* plants. The data represent means \pm SD ($n = 6$). **C**, Immunoblot analysis of photosynthetic proteins accumulated in mature and young leaves from 30-d-old wild-type and *sot5* plants. Coomassie Brilliant Blue (CBB) staining is shown to check the difference in sample loading. **D**, BN-PAGE analysis of photosynthetic complexes in thylakoid membranes isolated from 30-d-old wild-type and *sot5* plants. Each lane was loaded with 6 μ g of Chl.

true leaf of *sot5* had serrations and simplified vascular bundles (Fig. 3A), and cross sections of the first leaf showed that the epidermis of *sot5* was not as smooth as that of the wild type and that the air spaces between mesophyll cells were larger in *sot5* than in the wild type (Fig. 3A). In addition, the anatomical identity of palisade and spongy cells disappeared in *sot5* leaves (Fig. 3A), indicating that the leaf adaxial-abaxial polarity was altered in *sot5* mutants. Using scanning electron microscopy to examine the structure of abaxial and adaxial cells in *sot5* and wild-type leaves, we found that both adaxial and abaxial sides of *sot5* leaves were severely wrinkled and that adaxial epidermal cells resembled abaxial ones (Fig. 3B), suggesting reduced adaxial cell identity in the *sot5* mutant. Since *sot5* leaves are small and their mesophyll cells are loosely arranged, we hypothesized that the number of mesophyll cells was reduced in *sot5*. As expected, the

number of mesophyll cells in *sot5* was about 60% of the wild-type value (Fig. 3C), indicating that cell proliferation is inhibited in *sot5* leaves. Together, these results indicate that *SOT5* plays an important role in establishing abaxial-adaxial leaf polarity.

In Arabidopsis leaf development, ASYMMETRIC LEAVES1 (AS1) and AS2 form an epigenetic repressor complex that plays a critical role in repressing abaxial-determining genes and promotes adaxial polarity formation (Husbands et al., 2015). AS1 or AS2 loss-of-function mutants exhibit mild abaxialized leaves, such as lotus-like leaves (Xu et al., 2003). Previous work has demonstrated that the Arabidopsis mutants with altered leaf cell proliferation and leaf adaxial-abaxial patterning (*asymmetric leaves1/2 enhancer7*, *elongator*, and *scabra1* [*sca1*]) can enhance the *as2* phenotype, resulting in narrow and even needle-like (abaxialized) leaves (Yuan et al., 2010; Xu et al., 2012; Mateo-Bonmatí

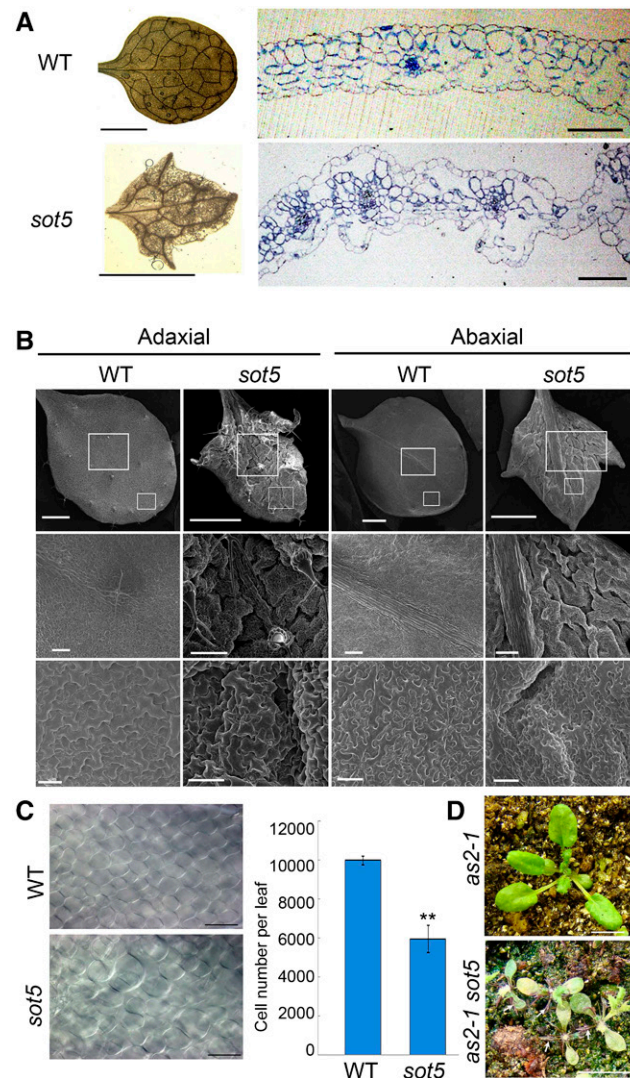


Figure 3. Leaf morphology and adaxial-abaxial polarity are altered in *sot5*. A, Morphology and cross section of the first true leaves from 25-d-old wild-type (WT) and *sot5* plants. Bars = 5 mm at left and 100 μ m at right. B, Scanning electron microscopy analysis of adaxial and abaxial surfaces of the first true leaves from 25-d-old wild-type and *sot5* plants. Enlargements of the larger and smaller squares in the top row are shown in the middle and bottom rows, respectively. Bars = 1 mm in the top row, 200 μ m in the middle row, and 50 μ m in the bottom row. C, Comparison of palisade cell size (left) and number (right) from the first true leaves of 25-d-old wild-type and *sot5* plants. The data represent means \pm SD of 10 leaves. Asterisks indicate a significant difference between the wild type and *sot5* (Student's *t* test, ** $P < 0.05$). Bars = 50 μ m. D, *sot5* enhances the leaf adaxial-abaxial polarity of *as2-1*. White arrows indicate filamentous leaves. Bars = 1 cm.

et al., 2015). To genetically test whether leaf polarity is altered in *sot5* mutants, we generated double *sot5 as2* mutants and found that they displayed many narrow or needle-like leaves (Fig. 3D), supporting our hypothesis that *sot5* is an enhancer of *as2*. Together, these results indicate that SOT5 has a role in promoting leaf adaxial identity formation.

SOT5 Encodes a Chloroplast-Localized PPR Protein

Genetic analysis showed that the *sot5* phenotype was controlled by a single nuclear recessive mutation.

To clone *sot5*, we produced an F2 population from the cross between *130-1* and *Landsberg erecta* (*Ler*) for map-based cloning. Using 900 recombinant plants, we mapped the *sot5* gene on the upper arm of chromosome 1 within an \sim 70-kb region between F26G16-a and T5I8-d markers, in which there are 20 annotated genes (Fig. 4A). Bioinformatic analysis revealed that, among these 20 candidates, six genes encode chloroplast-localized proteins, one of which is a PPR protein (AT1G30610), named EMB2279. DNA sequencing showed a G-to-A substitution at the first nucleotide of the seventh intron in AT1G30610 (Fig. 4B). Reverse

transcription (RT)-PCR analysis using the primer pair shown in Figure 4B detected three variants of the transcripts in *sot5* but only one in the wild type (Fig. 4C). Sequencing of these three transcripts revealed that the longest transcript (a in Fig. 4C) contained the seventh intron, the middle transcript (b in Fig. 4C) was the same as the wild type, and the shortest transcript (c in Fig. 4C) lacked 22-bp nucleotides at the 3' end of the seventh exon. The 22-bp nucleotide deletion resulted in a truncated form of SOT5. Thus, the *sot5* point mutation leads to a low efficiency of the intron splicing and missplicing of the exon. By contrast, we did not detect any alternative splicing events in the seventh intron of the *SOT5* gene in the wild type, which is contrary to recorded data in TAIR, where two transcriptional variants with or without the seventh intron of the *SOT5* gene are described. Based on our data, *SOT5* encodes only one protein with 978 amino acid residues.

To confirm the map-based cloning result, we transformed the *SOT5* cDNA under the control of the cauliflower mosaic virus 35S promoter into the *sot5* mutant. All the *sot5* phenotypes were complemented by over-expressing *SOT5* (Fig. 4D), confirming that *SOT5* encoding the 978-amino acid polypeptide is functional. A previous study reported that the null mutation *emb2279-2* is embryo lethal (Lurin et al., 2004; Cushing et al., 2005), causing embryo development to stop at the globular stage (Tzafirir et al., 2003). To determine whether *emb2279* and *sot5* are alleles of the same gene, we first identified *emb2279-2* (SALK_088420) from the Arabidopsis Biological Resource Center (ABRC) stock. Indeed, about 25% of the seeds were albino and aborted in the *emb2279-2* heterozygote siliques (Fig. 4E), confirming the embryo-lethal phenotype of *emb2279*. We then crossed *sot5* with the *emb2279-2* heterozygote. We found that half of the F1 plants were almost albino and half of them were wild type like when they were cultured on one-half-strength Murashige and Skoog medium. The albino plants could turn a bit green when growing on the medium but could not survive after transfer to the soil, since their leaves were severely pale (Fig. 4F). Furthermore, we generated *SOT5* knockdown lines by an artificial microRNA technique. Transgenic plants with low expression levels of *SOT5* showed a virescent phenotype (Fig. 4G). Taken together, our data suggest that *sot5* is a weak allele of *emb2279*.

Although *SOT5/EMB2279* was predicted to target chloroplasts, no evidence supports this prediction. To test this, we made a construct in which the first N-terminal 120 amino acids containing the transit peptide of *SOT5* were fused to yellow fluorescent protein (YFP) driven by the cauliflower mosaic virus 35S promoter and transformed it into protoplasts. Confocal microscopy analysis showed that YFP signal was totally overlapped with Chl autofluorescence (Fig. 5A), indicating that *SOT5* is localized in chloroplasts. RT-PCR analysis showed that *SOT5* was expressed ubiquitously in all examined tissues (Fig. 5B): in 5-d-old green or etiolated seedlings and in etiolated seedlings that had been exposed to light for 4 h (Fig. 5B). These results

indicate that *SOT5* functions in both green and non-green tissues.

SOT5 Is Required for Splicing of the Plastid *rpl2* and *trnK* Introns

SOT5 is a P-subfamily PPR protein composed of 978 amino acid residues. ScanProsite (<https://prosite.expasy.org/>) analysis showed that *SOT5* has 11 PPR motifs at the C terminus and no other function-known domains at the N terminus (Fig. 6A). BLAST searches demonstrated that *SOT5* is a single-copy gene in Arabidopsis and is conserved in monocots and dicots, but not in Bryophyta, Lycopodiophyta, Chlorophyta, and Rhodophyta (http://plants.ensembl.org/Arabidopsis_thaliana/Gene/Compar Ortholog?db=core;g=AT1G30610;r=1:10846513-10850724). The 11-PPR motifs of *SOT5* also are present in the mosses *Physcomitrella patens* and *Marchantia polymorpha*, in the fern *Selaginella moellendorffii*, but not in green algae (Supplemental Fig. S2).

Since PPR proteins participate in RNA processing, such as RNA splicing, editing, maturation, and stabilization, through directly recognizing specific RNA sequences, we propose that the target of *SOT5* also might be conserved in higher plants. To identify the targets of *SOT5*, we first predicted its binding target, the single-stranded RNAs, according to the PPR code and previously identified single-stranded RNAs targeted by different PPRs (Barkan et al., 2012; Yin et al., 2013; Yagi et al., 2014). Our analysis revealed that the 11 ribonucleotides targeted by the 11 PPR motifs of *SOT5* are XG(G/U)C(A/C)XXAUCX, with X representing any nucleotide that cannot be precisely predicted and nucleotides in parentheses being optional (Fig. 6B). We then searched the plastome using this potential *SOT5*-binding sequence and found that 14 plastid genes, ATCG00860 (*ycf2.1*), ATCG01280 (*ycf2.2*), ATCG00170 (*rpoC2*), ATCG00950 (*23S.1 rRNA*), ATCG01180 (*23S.2 rRNA*), ATCG00030 (*trnK*), ATCG01130 (*ycf1.2*), ATCG00100 (*trnG*), ATCG00360 (*ycf3*), ATCG00570 (*psbF*), ATCG00680 (*psbB*), ATCG00830.1 (*rpl2.1*), ATCG01310 (*rpl2.2*), and ATCG01100 (*ndhA*), contain the sequence (Supplemental Table S1). Based on these data, we designed primers flanking the binding sites and analyzed the expression levels of these candidate genes by reverse transcription-quantitative PCR (RT-qPCR). None of these genes were down-regulated significantly in *sot5* (Fig. 6C), but some NEP genes, such as *rpoC2*, *ycf1*, and *ycf2*, were up-regulated in *sot5* (Fig. 6C). These results indicate that *SOT5* is not required for the RNA stability of these candidate genes.

Among these candidate genes, *rpl2*, *ndhA*, *trnK*, and *trnG* contain the group II intron. Therefore, we investigated whether *SOT5* was involved in the intron splicing of these chloroplast genes. RT-qPCR analysis showed that the intron splicing efficiency for only *rpl2* was reduced significantly in *sot5* (Fig. 6D). Further RT-PCR analysis using the primers flanking the group II intron showed that the precursor levels of

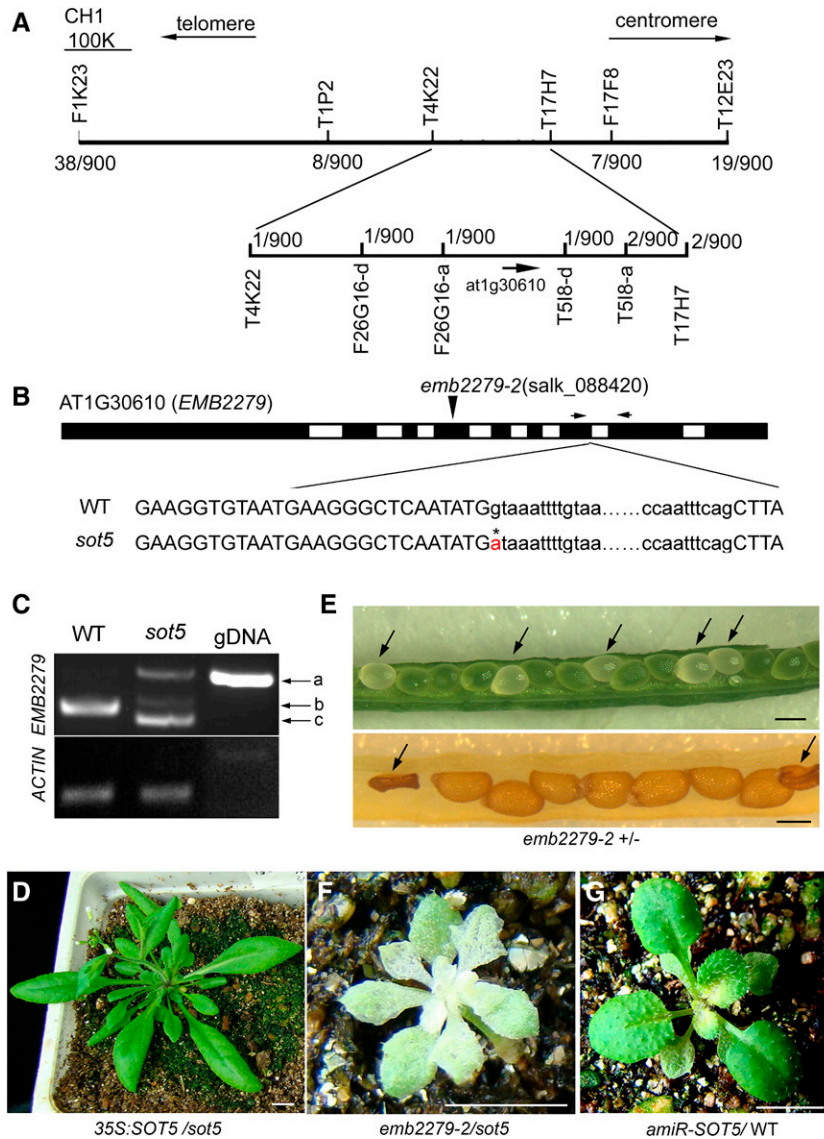


Figure 4. *SOT5* encodes *EMB2279*, a PPR protein. **A**, Map-based cloning of *SOT5*. *SOT5* was mapped between two markers, *F26G16-a* and *T518-d*, on chromosome 1. The molecular markers used for fine-mapping are shown. Numbers indicate the recombinant *sot5* mutants in 900 F2 individuals from crosses between *sot5* (Col-0 ecotype) and *Ler*. **B**, Model for the *SOT5* (AT1G30610) gene structure. The flanking sequences of the seventh intron are shown below the gene model. The G-to-A point mutation is labeled with an asterisk and red font in *sot5* at the first position of the seventh intron. The position of the T-DNA insertion in *emb2279-2* (SALK_088420) is shown above the gene model. The arrows show the primers used for RT-PCR. WT, Wild type. **C**, RT-PCR analysis of *SOT5* transcripts in Col-0 and *sot5* by using the primer pair shown in **B**. There are three bands, designated a, b, and c, in *sot5*. Genomic DNA (gDNA) was used as a temperate control, while the expression of *ACTIN* was used as a positive control. **D**, Overexpression of *SOT5* cDNA complemented the phenotype of *sot5*. Bar = 1 cm. **E**, The null mutation *emb2279-2* is embryo lethal. The top image shows the albino seeds (arrows) in a young *emb2279-2* heterozygote silique. The bottom image shows the aborted seeds (arrows) in an older *emb2279-2* heterozygote silique. Bars = 1 mm. **F**, Allelic test of *sot5* and *emb2279-2*. The F1 plant of *sot5/emb2279-2* appears albino and is seedling lethal. Bar = 1 cm. **G**, Knockdown lines of *SOT5* by the artificial microRNA technique exhibit a leaf virescent phenotype. Bar = 1 cm.

rpl2 transcripts were accumulated in *sot5* and *thf1* *sot5* but not in the wild type and *thf1*, but no difference was detected in transcript levels of other intron-containing genes between the wild type and *sot5* (Fig. 6E). Consistently, the levels of unspliced *rpl2* transcripts accumulated in the *SOT5* knockdown line

(*amiR-SOT5*) but were restored to wild-type levels in complementary lines (*35S::SOT5/sot5*; Fig. 6, F and G). Since mature tRNAs are small and have strong secondary structure and base modifications, they are difficult to detect precisely by RT-PCR (de Longevialle et al., 2008). To assess mature tRNA and precursor

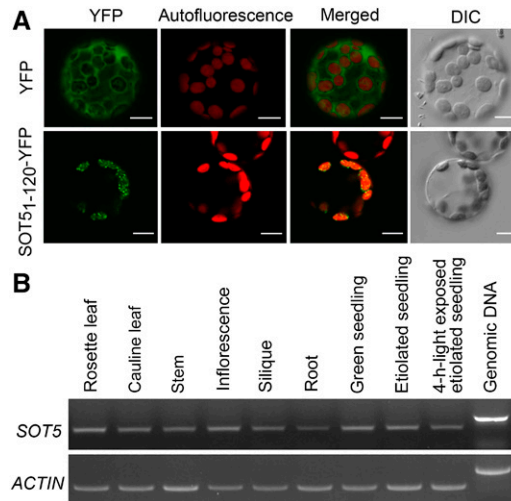


Figure 5. Subcellular localization of SOT5 and tissue expression patterns of *SOT5*. A, Microscopy analysis of the YFP fusion protein SOT5₁₋₁₂₀-YFP transiently expressed in Arabidopsis protoplasts. The green fluorescence of SOT5₁₋₁₂₀-YFP was overlapped with chloroplast autofluorescence in merged images. DIC, Differential interference contrast microscopy. Bars = 10 μ m. B, RT-PCR analysis of *SOT5* expression in various tissues and seedlings. Total RNA was extracted from rosette and cauline leaves, stems, inflorescences, siliques, and roots of 40-d-old plants. Total RNA was extracted from 5-d-old seedlings grown in light or darkness or etiolated seedlings exposed to light for 4 h. Genomic DNA was used as a template control, and *ACTIN* expression was used as a loading control of RNA.

levels, we employed northern-blot analysis. Our data demonstrated that only *trnK* unspliced precursors accumulated significantly in *sot5*, *sot5 thf1*, and *amiR-SOT5*, and *trnK* spliced mature products were reduced dramatically in these genotypes compared with the wild type and *35S:SOT5/sot5* (Fig. 6H), suggesting that *SOT5* mutations result in the reduced splicing efficiency of *trnK*. We were unable to verify the binding site of SOT5 in vitro, because the recombinant SOT5 protein was not expressed in prokaryotic and eukaryotic systems.

Since the *matK* gene, which is required for the splicing of seven group IIA introns in *trnA*, *trnI*, *trnV*, *atpF*, *rpl2*, *rps12-2*, and *trnK* (Zoschke et al., 2010), is located in the *trnK* intron, it is possible that lowered *trnK* intron splicing efficiency in *sot5* would affect the level of *matK* transcripts and subsequently affect the intron splicing of all group IIA genes. Moreover, decreased *rpl2* and *trnK* might affect plastid protein translation, leading to a decrease in MatK activity in *sot5*. However, our results demonstrated that the relative level of *matK* transcripts was enhanced in *sot5* (Fig. 6C), probably due to the higher accumulation of *trnK* precursors in *sot5*, and the splicing efficiency of *atpF*, *rps12-2*, *trnA*, *trnI*, and *trnV* was not affected significantly by *SOT5* mutations (Fig. 6, C and H). These results suggest that, in *sot5* mutants, MatK activity is high enough to maintain the splicing of these introns, and the reduced splicing efficiency of *rpl2* and *trnK* should be caused directly by *SOT5* mutations.

SOT5 Mutations Also Suppress the Leaf Variegation of *var2* and Inhibit Plastid rRNA Processing

Since *thf1* leaf variegation is attributed to a low level of *var2* (Zhang et al., 2009), we tested whether *sot5* can suppress the variegation phenotype of *var2*. Genetic analysis showed that *var2 sot5* double mutants displayed a similar phenotype to *sot5* mutants (Fig. 7A), suggesting that *sot5* is a suppressor of *var2*. This is consistent with the previously reported results that a decrease of plastid translation efficiency can suppress leaf variegation (Yu et al., 2008; Ma et al., 2015; Wu et al., 2016).

It was reported previously that mutants deficient in plastid translation display a reduced efficiency of plastid rRNA processing (Yu et al., 2008; Ma et al., 2015; Wu et al., 2016). Based on this, we investigated the effect of *sot5* mutations on rRNA processing. Northern-blot analysis showed that levels of mature 16S, 23S, and 4.5S rRNA were reduced in *sot5* compared with the wild type, whereas their precursors overaccumulated (Fig. 7B), similar to our previously reported mutants, *prp111-2* and *prps9* (Supplemental Fig. S3), which were defective in plastid ribosomal protein large subunit 11 and ribosomal protein small subunit 9, respectively (Wu et al., 2013; Ma et al., 2015). These results indicate that plastid rRNA processing is linked to ribosome biogenesis and that the *sot5* mutant is indeed defective in ribosome biogenesis and protein translation in plastids due to the reduced splicing of *rpl2* and *trnK* genes. The

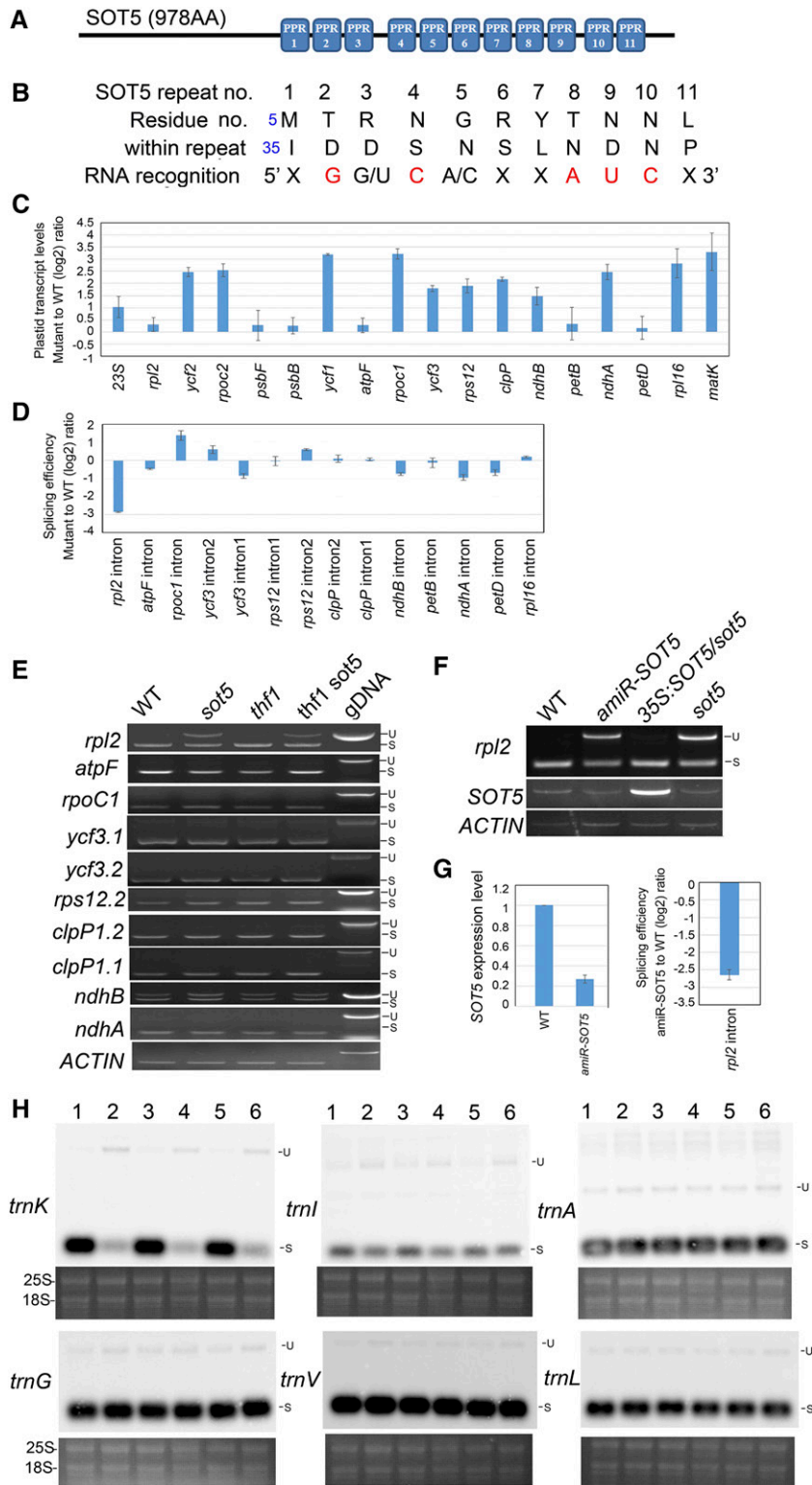


Figure 6. SOT5 is required for the splicing of the plastid *rp12* and *trnK* introns. A, Schematic SOT5 protein containing 11 PPR motifs. B, Predicted 11 ribonucleotides targeted by the 11 PPR motifs shown in A. In each repeat, the two key amino acid residues and their nucleotide targets are shown. X indicates unpredictable; two nucleotides with a slash mean optional; the red nucleotides are precisely predicted. C, RT-qPCR analysis of relative expression levels of the plastid genes containing the predicted target sequences or group IIA introns in *sot5*. Three biological replicates were analyzed. D, Splicing efficiency analysis of 14 plastid introns in *sot5* by RT-qPCR. Three biological replicates were analyzed. E, RT-PCR analysis of intron retention in eight plastid genes. Genomic DNA (gDNA) was used as a template control. S, Spliced; U, unspliced. F, RT-PCR analysis of *rp12*

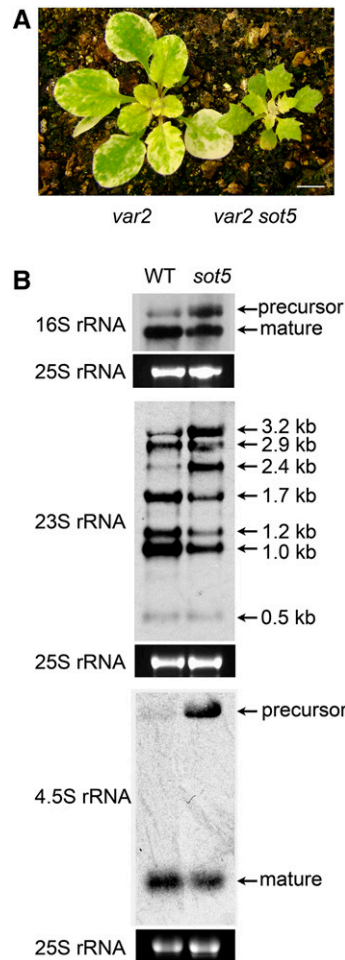


Figure 7. Loss-of-function *sot5* suppresses leaf variegation of *var2* and inhibits plastid rRNA processing. A, Phenotypes of 30-d-old *var2* and *var2 sot5* plants. *sot5* also can suppress the leaf variegation phenotype of *var2*. Bar = 1 cm. B, Northern-blot analysis of plastid rRNA processing in the wild type (WT) and *sot5*. Ethidium bromide staining of the 25S rRNA is shown below each blot to indicate differences in gel loading.

reduced efficiency of plastid rRNA processing may be the indirect effect of a translation defect in *sot5*. However, since 23S rRNA also was predicted to be the target of SOT5, we cannot rule out the possibility that SOT5 also can bind directly to 23S rRNA and facilitate its processing.

DISCUSSION

A Role of SOT5 in *rpl2* and *trnK* Intron Splicing

In this study, we identified a PPR protein, SOT5/EMB2279, that is involved in the splicing of the plas-

tid *rpl2* and *trnK* introns. Since both Rpl2 and trnK are essential for plastid protein translation, complete disruption of SOT5 would lead to embryo lethality in Arabidopsis, similar to previously reported results for EMB2279 (Lurin et al., 2004; Cushing et al., 2005). The *sot5* mutant contains a point mutation that leads to a low efficiency of the seventh intron splicing and missplicing of the seventh exon, resulting in a hypomorphic allele of *emb2279*. In TAIR, the SOT5 locus (AT1G30610) is recorded to produce two types of mRNA depending on whether the seventh intron is removed or not: the longer transcript encodes a 1,006-amino acid-long peptide with 10 PPR motifs,

Figure 6. (Continued.)

intron splicing in wild-type (WT), *amiR-SOT5/sot5*, *35S:SOT5/sot5*, and *sot5* plants. G, Analysis of SOT5 transcript levels and splicing efficiency of *rpl2* in wild-type and *amiR-SOT5* plants by RT-qPCR. H, Northern-blot analysis of plastid *trn* genes that contain introns in different genotypes. Lanes 1 to 6 indicate the wild type, *sot5*, *thf1*, *thf1 sot5*, *35S:SOT5/sot5*, and *amiR-SOT5*, respectively. The ethidium bromide-stained gel under each blot is shown as a loading control. Two biological replicates were analyzed, and one representative result is shown.

whereas the shorter one encodes a 978-amino acid-long peptide with 11 PPR motifs. However, we only detected the shorter transcript, which lacks 84 bp compared with the longer one, indicating that SOT5 has 11 PPR motifs. Consistent with the previous reports (Lurin et al., 2004; Cushing et al., 2005), total disruption of *SOT5* leads to embryo lethality in Arabidopsis, whereas knocking down *SOT5* expression through the artificial microRNA technique or partially knocking out the function of *SOT5* by a point mutation resulted in a virescent phenotype. These results are consistent with the fact that most of the suppressors for *thf1* and *var2* leaf variegation are mutants defective in the plastid translation system (Miura et al., 2007; Yu et al., 2008; Ma et al., 2015; Wu et al., 2016).

It is generally recognized that binding sites of PPR proteins that are involved in intron splicing are located in the intron (Schmitz-Linneweber et al., 2006; Beick et al., 2008; de Longevialle et al., 2008, 2010), which is helpful for maintaining RNA conformation and the subsequent recruitment of general splicing factors to facilitate intron splicing. Consistently, the predicted SOT5-binding site is located in the intron of *trnK*. In contrast, the predicted SOT5-binding site is located in the exon of *rpl2*, although there is a shorter motif containing eight nucleotides that overlapped with the predicted SOT5-binding site in the *rpl2* intron (523–530; Supplemental Table S1). Based on this, it is possible that SOT5 can facilitate both *rpl2* intron splicing and transcript stabilization. Considering that the plastid *rpl2* intron was independently lost in several species, such as in the Caryophyllales and Lythraceae (Downie et al., 1991; Gu et al., 2016), it will be interesting to examine whether the role of SOT5 homologs in these species is associated with *rpl2* transcript stabilization. In addition, we cannot exclude the possibility that the binding of SOT5 to the *rpl2* exon also is required for intron splicing. This hypothesis is consistent with the notion that the specific pattern of OTP51 action cannot be explained by an intron feature, because no conserved RNA motif was found in the *ycf3.2*, *atpF*, *trnV*, and *trnK* group IIA introns (de Longevialle et al., 2008). Whether PPR protein-binding sites can be located in an exon needs to be investigated further in the future.

Retrograde Signaling from Plastids to the Nucleus Regulates Leaf Development

Besides the leaf virescent phenotype, *sot5* displays multiple defects in leaf development, such as serrated leaves and less vascular bundles (Fig. 3). It is well documented that abnormal leaf morphology occurs frequently in mutants with dysfunctional chloroplasts: for example, in *DEFECTIVE CHLOROPLAST AND LEAVES*, functioning in plastid rRNA maturation (Keddie et al., 1996; Bellaoui et al., 2003); *SCA3*, encoding a plastidic RpoTp RNA polymerase (Hricová et al., 2006); *SCA1*, encoding a plastid-type ribosomal protein S5 (Mateo-Bonmatí et al., 2015); *ENF2*, encoding a chloroplast-localized and functionally unknown

protein, with low similarity to the bacterial polyamine transporters PotD and PotF (Tameshige et al., 2013); and *ANGULATA7*, encoding a DnaJ-like zinc finger domain protein localized in chloroplasts (Muñoz-Nortes et al., 2017). It has been suggested that the effect of chloroplast development on leaf morphology is associated with improper cell proliferation and expansion during leaf development (Mateo-Bonmatí et al., 2015). Recent studies showed that the serrated leaves and abnormal vascular development in *initiation factor3*, encoding a plastid translation initiation factor, were linked to altered auxin homeostasis and auxin-regulated pathways (Zheng et al., 2016); however, it remains unknown how chloroplast gene expression affects auxin biosynthesis and/or distribution. Our study also demonstrated that the layer of vertically elongated palisade cells beneath the adaxial epidermis was missing in *sot5* (Fig. 3), suggesting that adaxial domain formation is inhibited in *sot5* leaf primordia. Furthermore, the *as2 sot5* double mutant had more severe abaxialized filamentous leaves, which were similar to those of *sca1 as2* mutants. Both *sca1* and *sot5* are defective in a plastid ribosomal subunit (Mateo-Bonmatí et al., 2015), indicating that plastid translation may trigger a retrograde signal that regulates leaf polarity development. The *sot5* mutant provides ideal material in which to investigate the molecular mechanisms by which leaf development is feedback regulated by plastid gene expression for improved photosynthesis.

MATERIALS AND METHODS

Plant Materials and Growth Conditions

The Arabidopsis (*Arabidopsis thaliana*) ecotype Col-0 was used as the wild type in this study. Single mutants used in this study were described previously: *thf1-1* (Huang et al., 2006), *var2* (Yu et al., 2004) and *as2-1* (Xu et al., 2003). The *emb2279-2* (SALK_088420) mutant that was reported previously (Cushing et al., 2005) was obtained from the ABRC stock center (<https://abrc.osu.edu/>) and was genotyped according to the guidelines described there. Double mutants were identified in F2 generations derived from crosses between single mutants by the PCR-based genotyping procedure. Seeds were surface sterilized by 75% (v/v) ethanol and stratified at 4°C for 3 d and then sown onto one-half-strength Murashige and Skoog medium with 1% Suc, or seeds were sown directly into soil and grown in a phytotron with long-day conditions (16 h of light/8 h of dark) and light intensity (100 μmol photons m⁻² s⁻¹) at 22°C.

Map-Based Cloning, Plasmid Construction, and Transformation

Suppressors were identified by screening M2 seeds that were mutagenized by 0.1% (w/v) ethyl methanesulfonate in the *thf1* genetic background. The suppressor line was crossed to *Ler*, and the F2 seedlings were used as mapping populations. The mutation was mapped using simple sequence length polymorphism markers, and information is available at TAIR (<http://www.arabidopsis.org>). The *SOT5/EMB2279* coding DNA sequence (CDS) and the CDS for the first 120 amino acids including transit peptide sequence (fragment CDS) were cloned into the pENTR SD/D-TOPO entry vector (Invitrogen). The *SOT5/EMB2279* CDS was then recombined into the pGWB2 destination vector (Nakagawa et al., 2007). Artificial microRNA of *SOT5* was built according to Schwab et al. (2006). The destination vectors were transformed into Col-0 or *sot5* mutants according to the method described by Clough and Bent (1998). For transient subcellular localization analysis, the coding sequence of the first 120 amino acids including transit peptide sequence was recombined into the

p2GWY7 vector (<http://gateway.psb.ugent.be/vector/show/p2GWY7/search/index/>) and transformed into the Arabidopsis protoplasts according to the method of Wu et al. (2009).

Analyses of Chl Content, Chl Fluorescence, and Chloroplast Ultrastructure

Total Chl was extracted with 80% acetone at 4°C for 24 h in darkness. The levels of Chl *a* and Chl *b* were determined as described by Porra et al., 1989. Chl fluorescence was analyzed using Imaging PAM 101 (Walz), and F_v/F_m values were determined according to the manufacturer's instructions. For chloroplast ultrastructure analysis, the first true leaves of 15- and 30-d-old plants were sampled and fixed in a solution of 4% glutaraldehyde, then processed, embedded, and viewed via transmission electron microscopy as described (Zhou et al., 2009).

Protein Extraction, Immunoblot, and BN-PAGE Analyses

Protein extraction, immunoblot, and BN-PAGE analyses were performed as described previously (Huang et al., 2013).

Microscopy

Thin-section specimens and leaf samples for scanning electron microscopy were prepared according to our previous report (Huang et al., 2006). Samples for measuring and observing leaf mesophyll cells were processed as described by Xue et al. (2015).

RNA Isolation and RT-qPCR

Total RNA was extracted from leaves of 25-d-old plants using an RNeasy plant mini kit (Qiagen), according to the manufacturer's instructions. Downstream DNase I treatment and RT-PCR steps were conducted as described previously by Huang et al. (2013). Quantitative and semiquantitative RT-PCR of chloroplast genes was carried out using the gene-specific primers described previously by de Longevialle et al. (2008) and designed by ourselves (Supplemental Table S2). Three biological repeats were analyzed, and each sample was run in triplicate. The data set was normalized using *ACTIN2* as a reference. The method to quantify the transcript level and splicing efficiency of plastid genes was described previously (de Longevialle et al., 2008).

Northern-Blot Analysis

For northern-blot analysis of plastid tRNAs, the following biotinylated oligomer probes, which were described previously by de Longevialle et al. (2008), were synthesized by Thermo Fisher Scientific: *trnK*, 5'-AAAGC-CGAGTACTCTACCGTTG-3'; *trnL*, 5'-TGGGGATAGAGGGACTTGAAC-3'; *trnG*, 5'-CATCGTTAGCTTGGGAAGGCTAAGGG-3'; *trnI*, 5'-CTACCACT-GAGCTAATAGCCC-3'; *trnV*, 5'-GTGTAAACGAGGTGCTCTACCTAA-3'; and *trnA*, 5'-GGACTCGAACCCTGACATCCGCC-3'. Each probe sequence is in its exon. Total RNAs used for northern blot were extracted from 15-d-old seedlings according to the manufacturer's instructions (Promega Denaturing Solution; Z5651). A total of 1 to 2 µg of RNA was run on 2% agarose/formaldehyde gels, transferred to positively charged nylon membranes (Roche; 11417240001), and hybridized with biotinylated oligomer probes at 42°C. The North2South Chemiluminescent Hybridization and Detection Kit (Thermo Fisher Scientific; 17097) was used for the downstream steps according to the kit instructions. Northern-blot analysis of plastid rRNAs was performed as described previously (Wu et al., 2016).

Prediction of SOT5-Binding Sequences

SOT5-binding sequences were predicted as described previously (Barkan et al., 2012; Yin et al., 2013; Yagi et al., 2014; Shen et al., 2015, 2016). In brief, each PPR motif coordinates with one RNA base, and the combinations of the fifth and 35th amino acids (known as the PPR code) within each motif confer RNA-specific binding. The PPR motif sequences from SOT5 were used to search against the previously defined RNA recognition database for PPRs, leading to the following predicted recognition sequence: XG(G/U)C(A/C)XXAUCX, where X indicates a nucleotide that cannot be predicted precisely.

The Patmatch tool (<https://www.arabidopsis.org/cgi-bin/patmatch/nph-patmatch.pl>) was used to predict the binding sequences according to the SOT5 PPR code, XGKCMXXATCX (where X means any base, K means G or T, and M means A or C) by searching TAIR10 genes (+introns, +UTRs; DNA) database, with other parameters at default settings (Supplemental Table S1).

Accession Numbers

Sequence data from this article can be found in the TAIR data libraries under the following accession numbers: *EMB2279* (AT1G30610.2), *rpl2.1* (ATCG00830.1), *rpl2.2* (ATCG01310.1), *trnK* (ATCG00030.1), *ycf2.1* (ATCG00860.1), *ycf2.2* (ATCG01280.1), *rpoc2* (ATCG00170.1), *235.1* (ATCG00950.1), *235.2* (ATCG01180.1), *ycf1.2* (ATCG01130.1), *ycf3* (ATCG00360.1), *psbF* (ATCG00570.1), *psbB* (ATCG00680.1), *ndhA* (ATCG01100.1), *trnG* (ATCG00100.1), *trnA* (ATCG00940.1 and ATCG01190.1), *trnI* (ATCG00930.1 and ATCG01200.1), *trnV* (ATCG00450.1), and *trnL* (ATCG00400.1).

Supplemental Data

The following supplemental materials are available.

Supplemental Figure S1. Phenotypic comparison between wild-type and *sot5* seedlings.

Supplemental Figure S2. The PPR motifs of SOT5 are conserved in higher plants.

Supplemental Figure S3. Northern-blot analysis of plastid rRNA processing in *sot5*, *prpl11*, and *prps9* mutants.

Supplemental Table S1. Plastid genes that are predicted to be targets of SOT5.

Supplemental Table S2. Primer used in this study.

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