Functional Analysis of Polyprenyl Diphosphate Synthase Genes Involved in Plastoquinone and Ubiquinone Biosynthesis in Salvia miltiorrhiza

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INTRODUCTION

Plastoquinone (PQ) and ubiquinone (UQ, also known as Coenzyme Q, CoQ) are two major lipid-soluble prenylquinones involved in electron transfer and energy transformation in the plastids and mitochondria. PQs mainly present in higher plants, algae and cyanobacteria and are involved in photosynthesis, chlororespiration and carotenoid biosynthesis (Tian et al., 2007). UQs exist in all
aerobic organisms, such as plants and animals, and serve not only as coenzymes but also, in the reduced form, as antioxidants (Bentinger et al., 2007). Both PQs and UQs have the structural feature of a medium/long trans-polyisoprenyl side chain attached to the benzoquinone skeleton. Prenyl side chains are formed by sequential condensation of isopentenyl diphosphate (IPP) with allylic diphosphate in trans-configuration. The side chain of UQs contains variable numbers of isoprenoid units in different species. For instance, the side chain of UQ in Escherichia coli, Saccharomyces cerevisiae, and Schizosaccharomyces pombe contains eight (UQ-8), six (UQ-6), and ten isoprenoid units (UQ-10), respectively (Asai et al., 1994; Suzuki et al., 1997; Kainou et al., 2001; Ohara et al., 2010). In plants, the side chain of PQ usually contains nine isoprenoid units, whereas UQ side chain can be nine or ten isoprenoid units (Ducluzeau et al., 2012; Jones et al., 2013). For instance, Arabidopsis thaliana and most cereal crops contain UQ-9, whereas some vegetables, fruits and berries contain both UQ-9 and UQ-10 (Kamei et al., 1986; Mattila and Kumpulainen, 2001).

Although PQ and UQ play important roles in organisms, long-chain prenyl diphosphate synthase genes involved in the biosynthesis of PQ and UQ side chains have been functionally characterized only in A. thaliana, Oryza sativa, S. lycopersicum, and Hevea brasiliensis (Hirooka et al., 2003, 2005; Jun et al., 2004; Phatthiya et al., 2007; Ohara et al., 2010; Ducluzeau et al., 2012; Jones et al., 2013). In Arabidopsis, the main contributor that makes the solanesyl moiety of UQ-9 is At2g34630, which is a mitochondrion-located enzyme (Bouvier et al., 2000; van Schie et al., 2007; Danesi et al., 2011; Ducluzeau et al., 2012), whereas the genes encoding solanesyl diphosphate synthase targeted to chloroplasts and involved in PQ formation are At1g78510 and At1g17050 (Jun et al., 2004; Ducluzeau et al., 2012). Rice solanesyl diphosphate synthase genes involved in UQ-9 and PQ formation are OsSPS1 and OsSPS2, respectively. OsSPS1 was highly expressed in roots and localized in the mitochondria, whereas OsSPS2 was abundant in both leaves and roots and localized in the plastids (Ohara et al., 2010). S. lycopersicum solanesyl diphosphate synthase gene (SISPS) and decaprenyl diphosphate synthase gene (SIDS) are responsible for the production of PQ and UQ-10, respectively. SISPS was targeted to the plastids and involved in PQ biosynthesis, while SIDS could extend the prenyl chain length of UQ-6 to ten isoprenoid units and produce UQ-10 (Jones et al., 2013). In H. brasiliensis, HbSDS was involved in the synthesis of the prenyl side chain of PQ (Phatthiya et al., 2007). The Hevea solanesyl polyisoprenyl diphosphate synthase (PPS) gene involved in UQ formation has not been reported. Up to date, long-chain trans-type prenyltransferases genes related to PQ or UQ side chain biosynthesis in other higher plants remains unknown.

Salvia miltiorrhiza Bunge (Danshen in Chinese) is known for its great economic and medicinal value (Ma et al., 2012; Zhang and Lu, 2017). In recent years, it is emerging as a model medicinal plant for scientific research (Song et al., 2013; Deng and Lu, 2017). The genome of S. miltiorrhiza has been decoded (Zhang et al., 2015; Xu et al., 2016). In previous studies, we identified from the current S. miltiorrhiza genome assembly forty terpenoid biosynthesis-related genes, of which nine encode prenyltransferases (Ma et al., 2012). Based on the homologies of deduced amino acid sequences with other known prenyltransferases, the nine prenyltransferase genes identified from S. miltiorrhiza were predicted to encode short-chain prenyltransferases, including three geranylgeranyl diphosphate synthases (SmGGPPS1, SmGGPPS2, and SmGGPPS3), a farnesyl diphosphate synthase (SmFPPS), four heterodimeric geranyl diphosphate synthases (SmGPPS.LSU, SmGPPS.SSU1, SmGPPS.SSU1I.1, and SmGPPS.SSU1I.2), and a homodimeric geranyl diphosphate synthase (SmGPPS). To date, no S. miltiorrhiza genes encoding long-chain PPS have been characterized. Through genome sequence analysis and subsequent molecular cloning, we identified two long-chain prenyltransferases genes, termed SmPPS1 and SmPPS2, respectively. Comprehensive analysis showed that SmPPS1 were involved in PQ-9 biosynthesis, whereas SmPPS2 were involved in the biosynthesis of UQ-9 and UQ-10.

## MATERIALS AND METHODS

### Sequence Retrieval and Gene Identification

The amino acid sequences of plant solanesyl diphosphate synthases, including At1g17050, At1g78510 and At2g34630 in Arabidopsis, OsSPS1 and OsSPS2 in O. sativa and HbSDS in H. brasiliensis, were used for homologous search against the current assembly of S. miltiorrhiza (line 99–3) genome using tBLASTn algorithm in local server with an E-value cutoff of 1e-5 (Altschul et al., 1990, 1997; Xu et al., 2016). Exon/intron structures and coding sequences were predicted on the Genscan web server (Burge and Karlin, 1998) and corrected manually.

### Plant Materials and RNA Extraction

Plant materials used for tissue-specific expression and RACE experiments were 2-year-old, field-grown S. miltiorrhiza (line 99–3) plants. Flowers, leaves, stems, root cortices and root steles were collected and stored in liquid nitrogen until use. The plants used for analysis of light-induced expression of SmPPS1 and SmPPS2 were cultivated as described below. S. miltiorrhiza stem segments with nods and shoot tips were surface-sterilized and cultivated on MS agar media (Murashige and Skoog) at 25°C. The regenerated shoots were sub-cultured every 6 weeks. Four-week-old plantlets were transferred to soil and grown in a greenhouse for additional 4 weeks (18–30°C and about 14 h of light per day). Twenty uniformly growing shoots were selected for the analysis of light-induced expression of SmPPS1 and SmPPS2. Leaves were collected every 4 h from 16:00 pm on June 11th to 12:00 pm on June 12, 2013, when the time for sunrise was about 4:45 and for sunset was about 19:43. Plant materials were stored in liquid nitrogen immediately after collection.

Total RNA was extracted from plant tissues using the plant total RNA extraction kit (Bioteke, China). Genomic DNA contamination was eliminated by pretreatment of total RNA with RNase-free DNase (Promega, United States). RNA integrity was analyzed on 1% agarose gel. RNA quantity was determined

1. http://genes.mit.edu/GENSCAN.html
using a NanoDrop 2000C spectrophotometer (Thermo Fisher Scientific, United States).

Rapid Amplification of cDNA Ends and Cloning of Full Length SmPPS1 and SmPPS2

Total RNA isolated from roots of *S. miltiorrhiza* was used for mRNA purification using the Oligotex mRNA isolation kit (QIAGEN, Germany). First-strand cDNA was synthesized from the purified mRNA by SuperScript III reverse transcriptase (Invitrogen, United States) as per the manual. 5′ and 3′ RACE experiments were performed using the GeneRacer™ Kit (Invitrogen, United States). Gene-specific nested primers used for 5′ RACE of SmPPS1, respectively. Gene-specific nested primers used for 5′ and 3′ RACE of SmPPS1 were SmPPS1-R2 and SmPPS1-F3, respectively (Supplementary Table S1). Gene-specific primers used for 5′ and 3′ RACE of SmPPS2 are SmPPS2-R3 (5′ RACE nesting primer), SmPPS2-F2 (3′ RACE nesting primer), SmPPS2-R2 (5′ RACE nested primer), and SmPPS2-F3 (3′ RACE nested primer) (Supplementary Table S1). The predicted coding sequence of SmPPS1 was PCR-amplified using PPS1-F as the forward primer and PPS1-R as the reverse primer (Supplementary Table S2). The forward and reverse primers used for amplification of SmPPS1 coding sequence are PPS1-F and PPS1-R, respectively (Supplementary Table S2). Full length cDNA sequences of SmPPS1 and SmPPS2 were assembled using Vector NTI 7.1 (Lau et al., 2005).

Sequence Feature Analysis, Amino Acid Sequence Alignment and Phylogenetic Tree Construction

Sequence feature analysis was performed as described previously (Ma et al., 2012). The amino acid sequences of solanesyl diphosphate synthases from *Arabidopsis* (At2g34630, At1g17050 and At1g78510), rice (OsPPS1 and OsPPS2), *H. brasiliensis* (HbSDS), and *S. miltiorrhiza* (SmPPS1 and SmPPS2) were aligned using Clustal W algorithm-based AlignX, one of the tools of Vector NTI 7.1 software (Thompson et al., 1994; Lau et al., 2005), and then edited by GeneDoc software (Nicholas et al., 1997). For phylogenetic analysis, amino acid sequences of various known prenyltransferases were retrieved from GenBank. Plastid- and mitochondrion-targeting peptides were removed. The maximum likelihood tree was generated using the MABL tool2 with default parameters of the “one click” mode (Dereeper et al., 2008). The tree was displayed in the cladogram style with branch length ignored.

Quantitative Real-Time RT-PCR

Total RNAs isolated from various tissues of *S. miltiorrhiza* were reverse-transcribed into cDNAs using SuperScript III reverse transcriptase (Invitrogen, United States). Quantitative real-time RT-PCR was carried out in a 20 µL volume containing 6 µL diluted cDNA, 0.5 µM forward primer, 0.5 µM reverse primer and 1 × SYBR premix Ex Taq II (Tli RNaseH Plus) (TaKaRa Bio). PCR was performed using the Bio-Rad CFX96 system (Bio-Rad, Hercules, CA, United States) under the following conditions: predenaturation at 95°C for 30 s, annealing at 60°C for 30 s, extension at 72°C for 20 s, 39 cycles, and hold at 4°C. Three biological and three technical replicates were carried out. *SmUBQ10* was used as a reference gene as described previously (Ma et al., 2012). The forward and reverse primers used for SmPPS1 gene expression analysis are PPS1-F and PPS1-Rq, respectively (Supplementary Table S3). For SmPPS2, the forward and reverse primers are PPS2-Fq and PPS2-Rq, respectively (Supplementary Table S3). Relative abundance of transcripts was quantified using the comparative CT method as described previously (Livak and Schmittgen, 2008). Standard derivation was calculated from three independent biological replicates. SPSS version 21.0 was used to analyze the variance (ANOVA). *P* < 0.05 and *P* < 0.01 were considered as statistically significant and extremely significant, respectively.

Functional Complementation of Yeast coq1 Mutant

The open reading frames (ORFs) of SmPPS1 and SmPPS2 were inserted into the yeast expression vector pYES2-CT (Invitrogen, United States) using two restriction enzymes to give the final vectors pYES2-SmPPS1 (BamH I/Xba I) and pYES2-SmPPS2 (Kpn I/Not I). The empty vector pYES2-CT and the resulting constructs were separately introduced into the *S. cerevisiae coq1* mutant (strain 3138) using the yeast marker transformation kit (TaKaRa Bio, Japan). Functional complementation assay was performed as described previously (Ducluzeau et al., 2012). The yeast coq1 mutant with empty vector pYES2-CT and the wild type yeast strain BY4741 were used as controls.

Determination of UQ and PQ

Yeast cultivation, UQ induction and examination were carried out as described by Ducluzeau et al. (2012) with modifications. Three extractions were performed from each biological replicate. At least three independent biological replicates were carried out. Yeast cells were quantified by absorbance at 600 nm. Cells in 5 mL yeast solution were collected by centrifugation and then washed twice using 2 mL water. One milliliter water containing approximately 500 µL glass beads (0.5 mm) was added to the sediment and then vortexed for 10 min to destroy yeast cell wall. Samples were then quickly mixed with 2 mL acetone and extracted at room temperature for 30 min. The supernatant was evaporated to dryness in a vacuum desiccator at 40°C and re-suspended in 500 µL ethanol. For analysis of UQ in *S. miltiorrhiza*, leaves from 5-week-old *in vitro* cultivated plantlets were grinded into powder in liquid nitrogen and then extracted as described previously with slight modifications (Ohara et al., 2006). Leaf powder (100 mg) was quickly transferred into a 10 mL test tube with 5 mL ethanol, vortexed for 10 s, and then shaken at 120 rpm for 1 h at room temperature. After centrifuged at 10000 rpm for 15 min at 4°C, the supernatant was transferred to a new test tube and the solvent was evaporated in a vacuum desiccator at 40°C. The residue

2http://www.phylogeny.fr/
re-suspended in 500 µL of spectroscopically pure ethanol and then filtered through a Bond Elut-C18 solid phase extraction (SPE) column (Agilent, United States) and a 0.22 µm Millipore Express PES membrane filter (Merk Millipore, United States). The filtrate (10 µL) was injected into a C18 reverse-phase column (4.6 × 250 mm, Agilent, United States) and analyzed by HPLC-UV under the following conditions. Solvent system: ethanol/methanol (75:25); flow rate: 1.0 mL/min; temperature: 25°C; detection: absorbance measured at 275 nm. For analysis of PQ in S. miltiorrhiza, leaf powder was extracted with cold ethyl acetate as described by Kim et al. (2017) and Zbierzak et al. (2010). After centrifuged at 10000 rpm for 15 min at 4°C, UQ-6 (100 ng, Sigma) was added as internal standard (Zbierzak et al., 2010). The combined mixture was evaporated to dryness in a vacuum desiccator at 40°C and the residue was dissolved in 500 µL ethanol. HPLC-UV analysis was carried out using the isocratic solvent system consisting of ethanol/acetonitrile (1:3) at a flow rate of 1.5 mL/min for 26 min (Hunter et al., 2018). PQ-9 was quantified by comparison with the internal standard and detected spectrophotometrically at 255 nm (Zbierzak et al., 2010; Kim et al., 2017).

Subcellular Localization of SmPPS1 and SmPPS2

The ORFs of SmPPS1 and SmPPS2 without termination codon were PCR-amplified and inserted into pMD-18T (TaKaRa Bio, Japan) to yield T-SmPPS1 and T-SmPPS2 vectors. Sequence-verified clones were digested, gel-purified, and inserted into pCAMBIA1302-GFP, which had a cauliflower mosaic virus 35S promoter upstream of the cloning site. The resulting vectors, 1302-SmPPS1:GFP and 1302-SmPPS2:GFP (Figure 3A), were transformed into Agrobacterium tumefaciens strain EHA105 and then injected into leaf lamina of Nicotiana benthamiana plants according to the protocol described by Sparkes et al. (2006). After injection for 2–3 days, GFP fluorescence in N. benthamiana cells was observed under an Olympus FV3000 confocal microscope (Olympus, Japan) with an excitation filter of 488 nm from the argon laser. Wavelength of 488 nm was also used for fluorescence excitation of chlorophyll. MitoTracker Red CMXRos was used to stain the mitochondria with an excitation filter of 579 nm from the argon laser. Fluorescence emission was collected from 500 to 530 nm for GFP, 650 to 750 nm for chlorophyll, and 590 to 620 nm for MitoTracker Red CMXRos, respectively.

Binary Vectors Construction and Plant Transformation

The SmPPS1 and SmPPS2 coding sequences were cloned and inserted into the binary vector pCambia-1301 to make overexpression constructs. For knockdown of SmPPS1 and SmPPS2, two artificial microRNAs (amiRNAs), termed amiRPPS1 (5′-TTCTCACTGATGCGGATTGTC-3′) and amiRPPS2 (5′-TAGTACTCCATGCTGCAGCGA-3′), were designed as described by Schwab et al. (2006) and Shi et al. (2010). To construct knockdown vectors 1301-amiRPPS1 and 1301-amiRPPS2, P. trichocarpa ptc-miR408 and ptc-miR408* in a 0.9 kb cDNA of ptc-MIR408 gene were replaced with the designed amiRNA and amiRNA* through overlapping PCR. The resulting sequences were inserted into pCAMBIA1301 (Figures 5A, 7C). The constructs were introduced into A. tumefaciens strain GV3101 using the freeze-thaw method (Hofgen and Willmitzer, 1988). Agrobacterium-mediated transformation of S. miltiorrhiza was performed as described previously (Wang M. et al., 2017). Transgenic shoots were regenerated and selected on 1/2 Murashige-Skoog agar media supplemented with hygromycin 30 mg/L for about 20 days, and then excised and transferred to new 1/2 Murashige-Skoog agar media till rooting. Plantlets with well-developed leaves and roots were propagated. Transgenics were verified by PCR analysis of genomic DNA using HPTII gene specific primers, 5′-AGCTTGAACCTGCGCAGC-3′ and 5′-TGCCCTCGGAGTGACG-3′.

Chlorophyll, Carotenoid and Malondialdehyde (MDA) Content Analyses

Chlorophyll and carotenoids were extracted from the leaves of 5-week-old in vitro cultivated S. miltiorrhiza plants as described previously (Xing et al., 2010). Three extractions were carried out from each biological replicate. Light absorbance (A) of the final solution was measured at 663, 647, and 470 nm. The concentrations of chlorophyll a (Chla), chlorophyll b (Chlb) and carotenoids (Car) were calculated as described (Lichtenthaler, 1987). MDA content was determined using the thiobarbituric acid method with slight modifications (Dhindsa and Matowe, 1981). Leaves (0.2 g) from 5-week-old in vitro cultivated S. miltiorrhiza plants were homogenized in 3 mL 10% thiobarbituric acid and then stood at 4°C overnight. Supernatant was collected after centrifuged at 5000 rpm for 10 min at 4°C. Mixed 2 mL supernatant and 2 mL 0.6% thiobarbituric acid, immediately reacted in boiling water for 15 min, and then terminated by cooling down. The reaction mixture was centrifuged at 5000 rpm for 10 min at 4°C. Light absorbance (A) of the supernatant was measured at 450, 532, and 600 nm. The content of MDA was calculated as follows (Fan et al., 2012): MDA (µmol/L) = 6.456 × (OD532 − OD600) − 0.566 × OD450.

Transmission Electron Microscopy (TEM) and Scanning Electron Microscopy (SEM) Assays

Leaves harvested from 5-week-old wild type (WT) and amiRPPS1 transgenic plants were fixed in 2.5% glutaraldehyde for 1 week at 4°C. Tissue samples were rinsed three times in PBS buffer, soaked in 1% osmium tetroxide at room temperature for overnight, and dehydrated in a graded ethanol series. The samples were then embedded in Spurr resin for overnight and polymerized at 65°C for 2 days. Uranyl sections (70 nm) were cut with a Leica EM UC6 ultramicrotome (Leica, Germany), stained with 4.5% uranyl acetate for 5 min and lead citrate for 2 min, and then observed under a JEM-1400 TEM (JEOL, Japan).

Scanning electron microscopy analysis of leaves from 5-week-old WT and amiRPPS1 transgenic lines were carried out following the procedure of Irish and Sussex (1990) with some modifications. Leaf samples were fixed in 2.5% glutaraldehyde
overnight at 4°C, dehydrated in a graded ethanol series, and then immersed in tert-butanol for 30 min. The samples were dried in the JFD-320 Freeze Drying Device (JEOL, Japan), coated with gold dust in JFC-1600 auto fine coater (JEOL, Japan), and then placed on SEM stubs for examination under a JSM-6510LV SEM (JEOL, Japan).

RESULTS

Identification of Two PPS Genes in S. miltiorrhiza

Through BLAST analysis of previously known plant solanesyl diphosphate synthase sequences against the current assembly of S. miltiorrhiza (line 99-3) genome (Altschul et al., 1990, 1997; Xu et al., 2016), we identified two putative S. miltiorrhiza long-chain PPS genes, termed SmPPS1 and SmPPS2, respectively. In order to verify computational prediction and obtain full length cDNA sequences, 5’ RACE, 3’ RACE and RT-PCR were performed. The results showed that SmPPS1 had a 5’-untranslated region (UTR) of 210 bp, a 3’-UTR of 300 bp, and an ORF of 1266 bp. The length of 5’- and 3’-UTR of SmPPS2 was 155 and 125 bp, respectively, and its ORF was 1275 bp. BLAST analysis of SmPPS1 and SmPPS2 against the GenBank nucleotide database showed that SmPPS1 had the highest homology (80%) with H. brasiliensis HbSDS (Phatthiya et al., 2007). The deduced amino acid sequence of SmPPS1 showed 79% identity to Arabidopsis SPS isoform At1g78510. The results verify that SmPPS1 is a homolog of plant SPSs. The coding sequence of the cloned SmPPS2 cDNA showed four nucleotide differences with the predicted coding sequence of SmGPPS (GenBank accession number JN831107.1) (Ma et al., 2012). However, their amino acid sequences are identical. It suggests that SmPPS2 is full length cDNA of the previously predicted SmGPPS (Ma et al., 2012). Four nucleotide differences between SmPPS2 and SmGPPS may be caused by sequencing errors or single nucleotide polymorphisms. The deduced SmPPS2 protein sequence showed 70–83% identities with known SPSs in plant species, such as Sesamum indicum, Erythranthe guttata, Nicotiana attenuata, and 70–80% identities with homodimeric geranyl diphosphate synthases in various angiosperm plants, such as S. lycopersicum, Catharanthus roseus, Citrus sinensis, and Arabidopsis (Ma et al., 2012). The similarity between SPSs and S. miltiorrhiza, such as geranyl diphosphate synthases in various angiosperm plants, and 70–80% identities with homodimeric SPSs, such as At1g78510 and OsSPS2, whereas SmPPS2 has the same ASLHDVDL “FARM signature” with plastidial SPSs, such as At1g71050 and OsSPS2, whereas SmPPS2 has the same ASLHDVDL “FARM signature” with mitochondrial SPSs, such as At2g34630 and OsSPS1 (Figure 1B). Consistently, protein subcellular localization prediction on the TargetP 1.1 webserver3 showed that SmPPS1 was localized in the chloroplasts, whereas SmPPS2 was localized in the mitochondria. It suggests that SmPPS1 and SmPPS2 may be functionally distinct.

Phylogenetic analysis showed that plant trans-long-chain prenyltransferases could be divided into two different subgroups, including the plastid subgroup and the mitochondrion subgroup (Figure 1C). SmPPS1 belongs to the plastid subgroup, which includes PQ biosynthetic prenyltransferases, such as Arabidopsis At1g78510, At1g71050, rice OsSPS2, and other SPFs exclusively from oxygenic photosynthetic organisms (Figure 1C). SmPPS2 belongs to the mitochondrion subgroup, which includes UQ biosynthetic prenyltransferases, such as At2g34630, OsSPS1, and yeast COQ1 (Figure 1C) (Ohara et al., 2010; Ducluzeau et al., 2012). The results showed that SmPPS1 and SmPPS2 could be involved in the biosynthesis of PQ in the chloroplasts and UQ in the mitochondria, respectively.

Differential and Light-Induced Expression of SmPPS1 and SmPPS2

SmPPS2 has been previously shown to be ubiquitously expressed, although its expression in leaves and stems is slightly higher than in flowers, root cortices and root steles of 2-year-old, field-grown S. miltiorrhiza (Ma et al., 2012). In this study, we analyzed the expression of SmPPS1 in the same tissues used for the analysis of SmPPS2. The results showed that SmPPS1 was highly expressed in leaves, followed by stems (Figure 2A). The expression of SmPPS1 in flowers, root cortices and root steles was very low. It suggests

3http://www.cbs.dtu.dk/services/TargetP/
**FIGURE 1** | Comparative analysis of SmPPS1 and SmPPS2 in *S. miltiorrhiza* and various other plants. (A) Structure organization of SmPPS1 and SmPPS2 from *S. miltiorrhiza*, *Arabidopsis* and Rice. The exon/intron structures of coding regions were analyzed on the GSDS webserver (http://gsds.cbi.pku.edu.cn/) ([Hu et al., 2015]. Exons are represented by solid black boxes, introns are represented by lines between the boxes, and intron phases are indicated by 0, 1, and 2 ([Long et al., 1995]. GenBank accession numbers for full length SmPPS1 and SmPPS2 genes are JQ786411 and JN831107, respectively. The gene ID of OsSPS1 and OsSPS2 are AK071299 and AK066579, respectively. Different groups of SPSs are divided by dashed lines. (B) Amino acid sequence alignment of long-chain polyprenyl diphosphate synthases from various plant species. Seven domains conserved in eukaryotic polyprenyl diphosphate synthases are underlined and indicated by “I” to “VII”. The FARM signature and the SARM signature are indicated by “************” and “*****”, respectively. (C) Phylogenetic relationships of long-chain prenyltransferases from various organisms. The maximum likelihood tree was generated using the MABL tool (http://www.phylogeny.fr/). The accession numbers are listed as follows. SmPPS1, *S. miltiorrhiza* (MH924998); SmPPS2, *S. miltiorrhiza* (JX090100); At1g17050, *A. thaliana* (BT028962); At1g78510, *A. thaliana* (BT025560); At2g49830, *A. thaliana* (BT020524); HbSDS, *Hevea brasiliensis* (DQ437520); Cr-SPS, *Chlamydomonas reinhardtii* (XM_001693378.1); Cp-PT, *Cyanophora paradoxa* (NC_001675.1); ScCOQ1, *S. cerevisiae* (J05547); SpDPS1, *S. pombe* (D84311); Ec-ispB, *E. coli* (NP_417654.1); OsSPS1, *O. sativa* (AK071299); OsSPS2, *O. sativa* (AK066579).

**FIGURE 2** | Expression of SmPPS1 and SmPPS2 in *S. miltiorrhiza*. (A) Tissue-specific expression of SmPPS1 and SmPPS2 in flowers (Fl), leaves (Le), stems (St), root cortices (Rc), and root steles (Rs). Relative abundances of SmPPS1 and SmPPS2 in Rs and Fl are arbitrarily set to 1. (B) Different expression patterns of SmPPS1 and SmPPS2 in response to light induction in *S. miltiorrhiza* leaves. Light gray and white represent night and daytime, respectively.

SmPPS1 predominantly accumulates in green tissues. The results are consistent with the predicted functions of SmPPS1 in PQ biosynthesis in the chloroplasts and SmPPS2 in UQ biosynthesis in the mitochondria.

The expression patterns of SmPPS1 and SmPPS2 were analyzed on a summer day in Beijing. The sun rises at about 4:45 and sets at about 20:00, so the time for daytime and night is about 15 and 9 h, respectively. As a result, the transcription
of SmPPS1 and SmPPS2 were induced in day/night circulation with different expression patterns (Figure 2B). When daytime appeared, the transcript level of SmPPS1 began to accumulate rapidly and peaked at eight o’clock in the morning and then slowly decreased to a relatively stable level. The level of SmPPS2 decreased slightly during daytime and increased slowly at 20:00 when night began. SmPPS2 showed a relatively high expression level from midnight (zero o’clock) to four o’clock. Different expression patterns of SmPPS1 and SmPPS2 in response to light induction conformed to their predicated functions in the biosynthesis of PQ and UQ, respectively, which are two electron transporters in photophosphorylation and oxidative phosphorylation (Liu and Lu, 2016).

Subcellular Localization of SmPPS1 and SmPPS2

Prediction of protein subcellular localization using TargetP 1.1 showed that SmPPS1 and SmPPS2 could be localized in the chloroplasts and the mitochondria, respectively. In order to experimentally test the localization of SmPPS1 and SmPPS2 in planta, full length SmPPS1 and SmPPS2 were in-frame fused with a green fluorescent protein gene (GFP). The structures of the two resulting expression vectors, termed 1302-SmPPS1:GFP and 1302-SmPPS2:GFP, were shown in Figure 3A. Fusion genes were transformed into N. benthamiana leaves and transiently expressed. Observation of the transformed cells using a confocal laser scanning microscope showed that green fluorescence from SmPPS1:GFP completely coincided with red dots from chlorophyll auto-fluorescence (Figure 3B). It suggests that SmPPS1 was localized in the chloroplasts. Different from SmPPS1, SmPPS2:GFP showed a more complex localization. Most of the green fluorescence from SmPPS2:GFP showed a similar pattern to the Mitotracker red fluorescence in the mitochondria, whereas some of the other green fluorescence signals dissociated with the mitochondria. It indicated that SmPPS2 was, but not solely, localized in the mitochondria (Figure 3C).

Yeast Complementation of SmPPS1 and SmPPS2

It has been showed that the coq1 mutant of S. cerevisiae (3138) is unable to produce UQ and cannot grow on the minimum medium containing non-fermentable carbon source, since it lacks the mitochondrial hexaprenyl diphosphate synthase activity (Ashby and Edwards, 1996; Gin and Clarke, 2005). In order to further investigate the function of SmPPSs, SmPPS1 and SmPPS2 cDNAs were subcloned into the yeast D-galactose-inducible expression vector pYES2-CT to yield pYES2-SmPPS1 and pYES2-SmPPS2, respectively. The resulting constructs were introduced into the coq1 mutant 3138. As shown in Figure 4A,
the expression of SmPPS2 could restore the ability of mutant cells to use glycerol and ethanol as carbon sources, while no functional rescues were found for 3138 harboring pYES2-SmPPS1 or empty vector pYES2-CT. HPLC analysis showed that 3138 with pYES2-SmPPS2 produced UQ-6, UQ-9 and a small amount of UQ-10, and these UQs were not found in 3138 with pYES2-SmPPS1 or empty vector pYES2-CT (Figure 4B). The results indicate that SmPPS2 is a functional prenyltransferases involved in UQ-9 and UQ-10 production in S. miltiorrhiza. In addition to the peaks of UQ-6, UQ-9, and UQ-10, we noticed that there was a peak between the peaks of UQ-6 and UQ-9 in 3138 with pYES2-SmPPS2, and the peak in front of the peak of UQ-6 was changed significantly in different yeast cells analyzed. The compounds for these peaks remain to be identified.

**Functional Characterization of SmPPS1 in Transgenic S. miltiorrhiza Plants**

Further functional analysis of SmPPS1 was carried out using SmPPS1 transformed S. miltiorrhiza plants. The transgenics were obtained using the *Agrobacterium* -mediated transformation method (Wang M. et al., 2017). A total of four SmPPS1 overexpression lines, termed PPS1ox-4, PPS1ox-9, PPS1ox-23 and PPS1ox-29, respectively, were analyzed (Figure 5B). qRT-PCR showed that, among the four analyzed lines, PPS1ox-23 had the highest SmPPS1 expression level, which was about 9.4 times the level of SmPPS1 in WT plants (Figure 5C). Consistently, HPLC-UV analysis showed that PQ-9 content in PPS1ox-23 increased by 2.3 times compared with that in WT (Figure 5D). The lowest SmPPS1 expression level was detected in PPS1ox-4. It was about 2.3 times of the SmPPS1 expression level in WT (Figure 5D). PQ-9 content in PPS1ox-4 was 23% higher than that in WT (Figure 5D). The results suggest that SmPPS1 overexpression can increase PQ-9 content.

In order to down-regulate SmPPS1 expression in S. miltiorrhiza, the artificial microRNA method was used. Three SmPPS1 knockdown lines, named amiRPPS1-1, amiRPPS1-2 and amiRPPS1-8, respectively, were analyzed (Figure 5E). Among them, lines amiRPPS1-1 and amiRPPS1-2 showed significant decrease in SmPPS1 expression (Figure 5F). Consistently, PQ-9 content in these transgenic lines was reduced to below the detection limit (Figure 5G). AmiRPPS1-1 plants exhibited albino and dwarf phenotypes (Figures 6D,E), and leaf color might change to yellow during cultivation.
Further determination of chlorophyll and carotenoid contents showed that, compared with WT, the content of chlorophylls and carotenoids in amiRPPS1-1 leaves were dramatically reduced (Table 1). Chlorophyll a content decreased to 0.92% of WT, chlorophyll b reduced to 0.27%, carotenoids decreased to about 0.79%, and total chlorophylls decreased to about 0.13%. In addition, the ratio of chlorophyll a and chlorophyll b also decreased (Table 1). TEM observation of amiRPPS1-1 leaves showed that silence of SmPPS1 in S. miltiorrhiza resulted in abnormal development of chloroplasts. The chloroplasts in amiRPPS1-1 leaves lacked starch granules and thylakoids, and were smaller than the chloroplasts in WT (Figures 6F,G). SEM analysis showed that, compared to the WT, the development of trichomes was affected seriously in amiRPPS1-1 plants (Figures 6H,I). It seems that trichome development was blocked at very early developmental stage. AmiRPPS1-2, another SmPPS1 knockdown line with PQ-9 level below detection limit (Figure 5F), showed yellow and deep orange coloration on their leaves (Figure 6C). Since abnormal development of chloroplasts and color changes in amiRPPS1 transgenic lines, we speculated that the plants were undergoing oxidative stress. To test this hypothesis, the content of MDA, a reactive aldehyde and a biomarker of oxidative stress, was measured. The results showed significant increase of MDA content in amiRPPS1-1 and amiRPPS1-2 plants (Figure 6J), which confirmed the speculation. Taken together, knockdown of SmPPS1 gene may affect the biosynthesis of chlorophylls and carotenoids and the
FIGURE 6 | The phenotype and MDA content of amiRPPS1 transgenic *S. miltiorrhiza* lines. (A) wild-type plants. (B) amiRPPS1-1 plants. (C) amiRPPS1-2 plants. (D,E) WT (D) and amiRPPS1-1 (E) plants grown on 1/2 MS medium for 5 weeks. (F,G) TEM analysis of chloroplast development in leaf cells of WT (F) and amiRPPS1-1 (G). St, starch granule; TK, thylakoid. (H,I) SEM analysis of trichome development in WT (H) and amiRPPS1-1 (I). Arrows indicate trichomes. (J) MDA content in WT and amiRPPS1 lines. WT, wild type; 1, amiRPPS1-1; 2, amiRPPS1-2. ** means statistically extremely significant at *P* < 0.01.

| TABLE 1 | The content of chlorophylls and carotenoids in WT and amiRPPS1-1. |
|----------|-----------------|-----------------|-----------------|-----------------|-----------------|
|          | Chla (mg/g)     | Chlb (mg/g)     | Car (mg/g)      | Chl (mg/g)      | Chla/b          |
| WT       | 50 ± 5.1        | 14 ± 1.6        | 13 ± 1.2        | 64 ± 6.6        | 3.62 ± 0.11     |
| amiRPPS1-1 | 0.46 ± 0.23**  | 0.38 ± 0.25**  | 1.03 ± 0.3**   | 0.84 ± 0.3**    | 1.2 ± 0.17**    |

Leaves from 5-week-old plants were harvested for determination of chlorophyll and carotenoid contents. Differences of the means were compared between WT and amiRPPS1-1. Chla, chlorophyll a; Chlb, chlorophyll b; Car, carotenoid; Chl, total chlorophyll; Chla/b, chlorophyll a/chlorophyll b. *P* < 0.05 (*) and *P* < 0.01 (**) were considered statistically significant and extremely significant, respectively.
development of chloroplasts and trichomes, and cause toxic stress in cells.

**Determination of UQ in *S. miltiorrhiza* and Functional Characterization of *SmPPS2* in Transgenic Plants**

Yeast complementation assay showed that *SmPPS2* could functionally complement the *coq1* mutation and catalyzed the production of UQ-9 and UQ-10 in yeast cells (Figure 4). To verify the function of *SmPPS2* in *S. miltiorrhiza*, we analyzed the type and content of UQ in roots, stems and leaves of *S. miltiorrhiza* using HPLC-UV. The result showed that both UQ-9 and UQ-10 were produced in *S. miltiorrhiza* plants with the greatest content in leaves (27.97 ± 6.58 µg/g fresh weight (FW)), less in stems (8.85 ± 0.55 µg/g FW), and the least in roots (3.62 ± 1.13 µg/g FW) (Figures 7A,B and Table 2). The content of UQ-10 varies from 0.46 ± 0.29 µg/g FW in stems to 0.31 ± 0.04 µg/g FW in roots, which showed a gentle change compared with UQ-9 content varying significantly between 27.4 ± 6.58 µg/g FW in leaves and 3.32 ± 1.78 µg/g FW in roots. It is consistent with the results from yeast complementation assay.

To further analyze the function of *SmPPS2* in plants, transgenic *SmPPS2* overexpression and artificial miRNA lines were produced (Figures 7D,G). No visible phenotype changes were observed for these transgenic lines. Quantitative real-time PCR analysis of *SmPPS2* overexpression lines PPS2ox-5, PPS2ox-13, and PPS2ox-16 showed that the levels of *SmPPS2* transcripts were significantly up-regulated from 22 times of WT in PPS2ox-13 to 52.2 times in PPS2ox-16 (Figure 7E). Consistently, HPLC-UV analysis showed that total UQ contents were significantly increased in these *SmPPS2* overexpression lines (Figure 7F). The levels of UQ content in transgenics were from 152 to 163% of WT. It suggests that overexpression of *SmPPS2* can substantially increase UQ production in *S. miltiorrhiza* plants.

In artificial miRNA transgenic *S. miltiorrhiza* lines amiRPPS2-3 and amiRPPS2-6, the levels of *SmPPS2* transcripts were significantly down-regulated (Figure 7H). Consistently, total UQ content was decreased by 7% and 20% of WT in amiRPPS2-3 and amiRPPS2-6, respectively (Figure 7I). Reduction of *SmPPS2* transcripts and total UQ content were also detected in amiRPPS2-22, although the reduction was not statistically significant (Figures 7H,I). These data suggest that alteration of *SmPPS2* expression can affect the accumulation of UQ in *S. miltiorrhiza*.

**DISCUSSION**

*Salvia miltiorrhiza* is a widely used traditional Chinese medicine material and an emerging model system for medicinal plant biology (Ma et al., 2012; Song et al., 2013; Deng and Lu, 2017; Zhang and Lu, 2017). So far, more than two hundred chemical compounds have been isolated from this plant (Wang L. et al., 2017; Mei et al., 2019). Lipophilic diterpenoid tanshinones and hydrophilic phenolic acids are two major classes of bioactive compounds in *S. miltiorrhiza* (Gao et al., 2009; Kai et al., 2010; Hao et al., 2013; Xing et al., 2018). Other compounds, such as quinones, flavonoids, anthocyanidins, alkaloids, monoterpenes, sesquiterpenes, triterpenes, sterols and saccharides, were also found in *S. miltiorrhiza* (Wang L. et al., 2017; Mei et al., 2019), although their medicinal value is largely unknown.

Plastoquinone and UQ are two major classes of prenylquinones. In plants, PQ usually contains nine isoprenoid units, whereas the number of isoprenoid units for UQ can be nine or ten, of which UQ-10 is clinically prescribed as a complementary medicine in treating cardiovascular diseases, such as hypertension, hyperlipidemia, coronary artery disease and heart failure (Liu and Lu, 2016). UQ-10 supplementation energizes the body and increases body energy production in the form of ATP. It is recognized as a kind of super vitamin and known as vitamin Q (Shukla and Dubey, 2018). Through HPLC-UV analysis, we reported for the first time that both UQ-9 and UQ-10 were produced in *S. miltiorrhiza*. Although the medicinal significance of UQ-10 in this plant remained to be elucidated, it is possible that UQ-10 may strengthen the therapeutic effect of *S. miltiorrhiza* in clinical practice.

It has been shown that the content of UQ varies significantly in different organs of a plant and in different plant species. In *O. sativa*, UQ-9 content is 8.7 ± 5.0, 9.3 ± 3.7, and 7.7 ± 2.0 µg/g dry weight (DW) in roots, stems and leaves, respectively, whereas it is only 1.4 ± 0.2 µg/g DW in grains (Ohara et al., 2006). Among vegetable oils, maize germ oil and sunflower refined oil contain abundant UQ-9. The content of UQ-9 in maize germ oil and sunflower refined oil is 233.71 and 101.3 µg/g, respectively (Rodriguez-Acuña et al., 2008; Pyo, 2010). In soybean and rapeseed oils, UQ-10 contents are relatively high, which are 92.3 µg/g (Kamei et al., 1986) and 63.5 µg/g (Mattila and Kumpulainen, 2001), respectively. In addition, some other vegetables and fruits also contain UQ-9 and UQ-10. However, the contents are much less compared with those in vegetable oils. For instance, the content of UQ-9 and UQ-10 in pea, cauliflower, apple and strawberry are 0.1 and 1.7 µg/g FW, 0.04 and 2.7 µg/g FW, 0.2 and 1.3 µg/g FW, and 0.1 and 1.4 µg/g FW, respectively (Mattila and Kumpulainen, 2001). In this study, we found that both UQ-9 and UQ-10 were produced in leaves, stems and roots of *S. miltiorrhiza* plants. The highest total UQ content is 27.97 ± 6.58 µg/g FW, which was found in leaves. Generally speaking, the content of total UQ in *S. miltiorrhiza* is higher than that in rice, vegetables and fruits but less than that in vegetable oils (Kamei et al., 1986; Mattila and Kumpulainen, 2001; Ohara et al., 2006; Rodriguez-Acuña et al., 2008; Pyo, 2010).

The pathway and enzyme genes of UQ biosynthesis have been well studied in bacteria and yeasts; however, they are largely unknown in plants (Liu and Lu, 2016). For instance, the current model for UQ biosynthesis in plants was proposed based on the biosynthetic pathway of yeast UQ and has not been experimental demonstrated (Hayashi et al., 2014); the majority of UQ biosynthetic enzyme genes have not been functionally analyzed in plants, although orthologs have been identified from *Arabidopsis* and are able to complement yeast mutants; *Arabidopsis* does not contain the ortholog of *Coq7*, which encodes a mono-oxygenase involved in the penultimate step of UQ biosynthesis (Hayashi et al., 2014); in addition, a few
FIGURE 7 | Analysis of UQ in wild-type and SmPPS2 overexpression and knockdown S. miltiorrhiza plantlets. (A) Separation of UQ-9 and UQ-10 standards; (B) Analysis of UQ-9 and UQ-10 in S. miltiorrhiza leaves. (C) Schematic representation of T-DNA in 1301-SmPPS2 and 1301-amiRPPS2 plasmids. LB, left boundary of T-DNA; Poly, poly(A) signal; Hyg, hygromycin phosphotransferase gene; 35S, the promoter of CaMV 35S; GFP, green fluorescent protein gene; Nos, the terminator of nopaline synthase gene; RB, right boundary of T-DNA. (D) PCR verification of WT and SmPPS2 overexpression lines. (E) Relative expression level of SmPPS2 in WT and SmPPS2 overexpression lines. (F) UQ content in WT and SmPPS2 overexpression lines. (G) PCR verification of WT and SmPPS2 knockdown lines. (H) Relative expression level of SmPPS2 in WT and SmPPS2 knockdown lines. (I) UQ content in WT and SmPPS2 knockdown lines. WT, wild type; 5, 13 and 16, SmPPS2 overexpression lines PPS2ox-5, PPS2ox-13 and PPS2ox-16; 3, 6 and 22, SmPPS2 knockdown lines amiRPPS2-3, amiRPPS2-6 and amiRPPS2-22. $P < 0.05$ (*) and $P < 0.01$ (***) were considered statistically significant and extremely significant, respectively.
UQ biosynthetic enzyme genes are still unknown even in yeast cells (Hayashi et al., 2014). Moreover, although PQ biosynthetic pathway is relatively clear, genes involved in PQ biosynthetic pathway have mainly studied in Arabidopsis.

Polypropenyl diphosphate synthases are key enzymes to determine the number of isoprenoid units in the biosynthesis of isoprenoid compounds and their derivatives, such as monoterpens, diterpenes, triterpenoids, PQ, UQ, and vitamin E (Liu and Lu, 2016). Based on the chain length of final products, PPSs can be divided into three classes, including short-, medium- and long-chain PPSs (Hemmi et al., 2002). Among them, long-chain PPSs are significant to PQ and UQ biosynthesis in plants. So far, PPS genes have been identified from Arabidopsis, rice, tomato and H. brasiliensis (Jun et al., 2004; Phaththiya et al., 2007; Ohara et al., 2010; Ducluzeau et al., 2012; Jones et al., 2013). However, functional characterization through transgenic plants has only been performed for Arabidopsis At2g34630 and tomato SlSPS and SlDPS (Ducluzeau et al., 2012; Jones et al., 2013). Analysis of Arabidopsis At2g34630 T-DNA mutants and overexpressing lines showed that At2g34630 was involved in UQ-9 biosynthesis (Ducluzeau et al., 2012). Through virus-induced gene silencing of SlSPS and SlDPS and overexpression of tomato SlSIPS in tobacco, Jones et al. (2013) showed that tomato SlDPS is involved in UQ-10, whereas SlSPS is involved in PQ-9 biosynthesis. In this study, we identified and characterized two long-chain SmPPS genes from S. miltiorrhiza. The cloned nucleotide sequence of SmPPS2 is almost identical to the ORF of SmGPPS previously annotated to be involved in terpenoid biosynthesis in S. miltiorrhiza (Ma et al., 2012), and the deduced amino acid sequence of SmPPS2 and SmGPPS are identical. It suggests that the cloned SmPPS2 is full length cDNA of the annotated SmGPPS. Similar case of SmPPS2 was found for Arabidopsis At2g34630 (AtPPPS), which had been annotated as AtGPPS, but later was proved to be a SP involved in the biosynthesis of UQ side chain (Hsieh et al., 2011; Ducluzeau et al., 2012). The reason causing erroneous annotation is probably that GPPSs are evolutionarily most closely related to long-chain PPSs in angiosperms (Hsieh et al., 2011). Functional analysis of SmPPSs through stable gene overexpression and silencing in the original plant species showed that SmPPS1, similar to SlSIPS, was involved in PQ-9 biosynthesis, whereas SmPPS2 was involved in the biosynthesis of UQ-9 and UQ-10, with the major product to be UQ-9. It suggests the difference among the function of SmPPS2, SlDPS, and At2g34630.

In consistent with their functions in photophosphorylation-related PQ and oxidative phosphorylation-related UQ, SmPPS1 and SmPPS2 showed differential responses to light induction. When a day begins, plant photosynthesis and photophosphorylation are enhanced with the increase of sunlight. Up-regulation of SmPPS1 may facilitate PQ production to help the photophosphorylation process. Oxidative phosphorylation in respiration has little relevance to light, so the expression of SmPPS2 was relatively stable during daytime.

Experimental GFP analysis of N. benthamiana leaves showed that the green pseudocolor of SmPPS1-GFP was co-localized with the red pseudocolor of chloroplast autofluorescence. It suggests that SmPPS1 is localized in the chloroplasts. The results are consistent with previous results for At1g78510, OsSPS2 and SlSIPS involved in PQ side chain biosynthesis (Ohara et al., 2010; Ducluzeau et al., 2012; Block et al., 2013; Jones et al., 2013). Experimental analysis of SmPPS2-GFP localization showed that the green fluorescence was not only co-localized with the red fluorescence of mitochondria stained with a marker dye MitoTracker, but also found outside of the mitochondria. It suggests that, in addition to the mitochondria, SmPPS2 is probably localized in other organelles. Similarly, it has been shown that At2g34630, which is involved in UQ side chain biosynthesis in Arabidopsis, targets to both mitochondria and chloroplasts (Ducluzeau et al., 2012). Tomato SIDPS:GFP showed diffused fluorescence in N. benthamiana cells (Jones et al., 2013). The results are consistent with the report that solanesyl moiety was attached to benzoquinone through the Golgi/ER system in spinach leaves (Swiezewska et al., 1993). It indicates that the intermediates of UQ biosynthesis may move from one organelle to another in a cell (Ohara et al., 2004). Moreover, distinct function of SmPPS1 and SmPPS2 in coq1 could be due to different subcellular localization (Jun et al., 2004; Ohara et al., 2010). For instance, adding a mitochondrial signal peptide to the N-terminus of Arabidopsis At1g78510 could result in the complementation of S. pombe dp1 and dps1 mutant (Jun et al., 2004).

SmPPS1 down-regulated S. miltiorrhiza plants with exhibited abnormal chloroplast development, abnormal trichome development, and varied leaf bleaching phenotypes. The phenotypes are similar to that found in SIPS-silenced tomato transgenics (Jones et al., 2013). PQ is a cofactor of phytoene desaturation and is indirectly involved in the biosynthesis of carotenoids, which, together with PQ, are important antioxidants in protection of cell membranes against oxidative stress (Norris et al., 1995; Carol and Kuntz, 2001). Plants lacking PQ and carotenoids will be vulnerable to photo-oxidative damage and develop into photo-bleached seedlings (Chalker-Scott, 1999; Gould, 2004; Ksas et al., 2015; Liu and Lu, 2016). Abnormal chloroplast development in SmPPS1 down-regulated plants could also be caused by photo-oxidative stress, since significant increasing of MDA content was observed. Homogentisate solanesyltransferase (HST) is the other key enzyme involved in PQ biosynthesis. Mutation of HST may also result in PQ

| Tissue | UQ content (µg/g FW) | Total UQ |
|--------|-----------------------|----------|
| Root   | 3.32 ± 1.78           | 3.62 ± 1.13 |
| Stem   | 8.39 ± 0.52           | 8.85 ± 0.55 |
| Leaf   | 27.4 ± 6.58           | 27.97 ± 6.58 |

The means ± SD of three replicates are shown. FW, fresh weight.

**TABLE 2 | UQ content in roots, stems and leaves of S. miltiorrhiza.**

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deficiency and albinism phenotype, such as Arabidopsis pds2 mutant that has a 6bp-lesion in AtHST gene (Tian et al., 2007). However, pds2 mutant showed stomata closure defects, which was not found in SmPPS2 knockdown S. miltiorrhiza plants. It suggests the difference between PPS and HST down-regulated plants.

No visible phenotype changes were observed for SmPPS2 transgenics, but down-regulation of SmPPS2 might result in decrease of total UQ content, and up-regulation of SmPPS2 might cause UQ accumulation. It suggests that SmPPS2 is involved in UQ biosynthesis. The levels of SmPPS2 transcripts in SmPPS2 overexpression lines were significantly up-regulated from 22 to 52.5 times of WT. However, the content of total UQ was increased only by 52–63%. Similar phenomenon was observed for overexpression of At2g34630 in Arabidopsis (Ducluzeau et al., 2012). It indicates that overexpression of long-chain PPS gene has limited effects on plant UQ production. This is because long-chain PPS, which provides the prenyl side chain precursors for UQ structure, is not the rate-limiting enzyme of UQ biosynthesis (Liu and Lu, 2016). In contrast, overexpression of the rate-limiting enzyme gene PPT/COQ2 could effectively improve UQ production in plants (Ohara et al., 2004; Stiff, 2010). From a biotechnological point of view, comprehensive approaches, such as simultaneously increasing side chain supply and activating rate-limiting steps, could be efficient ways to raise UQ content in transgenic plants (Kumar et al., 2012; Liu and Lu, 2016).

Taken together, this is the first report for UQ biosynthesis in a valuable medicinal model plant. The results provide useful information for further elucidation of PQ and UQ biosynthetic pathways and for the production of clinically significant UQ-10 through metabolic engineering and synthetic biology approaches.

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DATA AVAILABILITY

The datasets generated for this study can be found in NCBI GenBank, MH924998, JX090100.

AUTHOR CONTRIBUTIONS

SL conceived and designed the experiments. ML, YM, QD, XH, and MW performed the experiments. ML, YM, and SL wrote the manuscript. All authors approved the final version of the manuscript.

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SUPPLEMENTARY MATERIAL

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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