Overexpression of SmSCR1 Promotes Tanshinone Accumulation and Hairy Root Growth in Salvia miltiorrhiza

Wei Zhou1†, Shuai Wang1†, Yafang Shen1†, Yunhui Liu1, Itay Maoz2, Xiankui Gao1, Chengan Chen1, Tingyao Liu1, Can Wang1 and Guoyin Kai1*

1 Laboratory for Core Technology of Traditional Chinese Medicine (TCM) Quality Improvement and Transformation, School of Pharmaceutical Sciences, The Third Affiliated Hospital, School of Pharmacy and Academy of Chinese Medical Science, Zhejiang Chinese Medical University, Hangzhou, China, 2 Department of Postharvest Science, Agricultural Research Organization, Volcani Center, Rehovot, Israel

Lipid-soluble tanshinone is one of the main bioactive substances in the medicinal plant Salvia miltiorrhiza, and its medicinal demand is growing rapidly. Yeast extract (YE) modulates the tanshinone biosynthesis, but the underlying regulatory network remains obscure. In this study, a YE-responsive transcriptional factor Scarecrow1 (SCR1) was identified in S. miltiorrhiza from the YE-induced transcriptome dataset. SmSCR1 is located in the nucleus. Overexpression of SmSCR1 in S. miltiorrhiza roots resulted in a significantly higher accumulation of tanshinone than the control, with the highest 1.49-fold increase. We also detected upregulation of tanshinone biosynthetic genes, SmSCR1 and SmHMGR1, and distinct alteration of growth and development of the hairy roots in the overexpression lines compared to the control. An inverse phenotype was observed in SmSCR1-SRDX suppression expression lines. We found that SmSCR1 can bind to the promoter of SmCPS1 to induce its expression. This study provides new insight into the regulatory mechanism on the growth and development of hairy roots, tanshinone accumulation, and the metabolic engineering of bioactive compounds in S. miltiorrhiza.

Keywords: Salvia miltiorrhiza, GRAS transcription factor, tanshinone accumulation, hairy root growth, regulatory mechanism

INTRODUCTION

Salvia miltiorrhiza Bunge is a Chinese herbal medicine that belongs to the Salvia genus in the Lamiaceae family (Jia et al., 2019; Jung et al., 2020). In Asian countries, dried roots and stems of S. miltiorrhiza are a common treatment for cardiovascular system-related diseases (Hao et al., 2015). The bioactivity of S. miltiorrhiza is associated with lipid-soluble tanshinones, including tanshinone I (TA-I), cryptotanshinone (CT), dihydrotanshinone (DT), and tanshinone IIA (TA-II) (Fu et al., 2020). Increasing attention has been given to the abovementioned compounds in recent mechanism, mainly focusing on improving the yield of tanshinone in S. miltiorrhiza.

Abbreviations: CPS1, copalyl diphosphate synthase I; HMGR1, 3-hydroxy-3-methylglutaryl-CoA-reductase I; GGPPS1, geranylgeranyl diphosphate synthase I; HMGs, hydroxymethylglutaryl-CoA synthase; IPPI, isopentenyl diphosphate isomerase; KSL, ent-kaurene synthase like; CPS1, copalyl diphosphate synthase I; CYP76AH1, miltiradiene oxidase; DXR, 1-deoxy-D-xylulose 5-phosphate reductoisomerase; DXS, deoxy-D-xylulose 5-phosphate synthase synthase II; Dual-LUC, dual-luciferase; YE, yeast extract; GRASI, GRAS transcriptional factor I; DT, dihydrotanshinone; CT, cryptotanshinone; TA-I, tanshinone I; TA-II, tanshinone IIA; TTA, total tanshinone.
Tanshinones are a type of diterpenoids produced in two phases in *S. miltiorrhiza* roots (Supplementary Figure 1). First, common terpenoid precursors [e.g., isopentenyl diphosphate (IPP) and dimethylallyl diphosphate (DMAPP)] are generated by two distinct processes (e.g., mevalonate (MVA) pathway in cytosol and 2-C-Methyl-D-erythritol-4-phosphate (MEP) pathway in plastids) (Ma et al., 2012). The universal diterpenoid precursor, geranylgeranyl diphosphate (GGPP), is then biosynthesized GGPP synthase (GGPPS) (Shi et al., 2016a,b). Then, three known synthases, namesely, copalyl diphosphate synthase 1 (CPS1), kaurene synthase-like 1 (KSL1), and mitilradiene oxidase (CYP76AH1), and a still-unidentified enzyme(s) produce tanshinones (Gao et al., 2009; Xing et al., 2018). Overexpression or downregulation of one or two synthases significantly impacts tanshinone synthesis (Kai et al., 2011; Wang et al., 2015; Ma et al., 2016). Plant transcription factors (TFs) play important regulatory roles in resistance to stress and metabolic engineering. TFs can regulate one or multiple biosynthetic genes from a single or multiple pathways (Xu et al., 2016; Wang et al., 2017; Yang et al., 2017). TFs are a diverse group of genes including many families and subfamilies, such as bHLH, MYB, ERF, bZIP, and GRAS (Deng et al., 2020a,b; Hao et al., 2020; Shi et al., 2021a,b; Zhou et al., 2021a,b). Less is known about the regulatory actions or target genes of the GRAS TFs, which play a critical role in the tanshinone biosynthetic pathway in *S. miltiorrhiza*.

The GRAS TF family has been discovered to be unique in plants. GRAS TFs consist of three gene members, namely, gibberellic acid insensitive (GA1), repressor of GA1-3 mutant (RGA), and Scarecrow (SCR). A whole-genome analysis from *Arabidopsis thaliana*, rice (*Oryza sativa*), and mus lily (*Lilium longiflorum*) grouped the GRAS families into ten distinctive subfamilies, namely, SCR, short-root (SHR), *L. longiflorum* SCR-like (LISCL), DELLA, SCL-like 3 (SCL3), SCL-like 4 and SCL-like 7 (SCLA/7), *A. thaliana* Lateral suppressor (AtLAS), hairy meristem (HAM), *A. thaliana* phytochrome A signal transduction 1 (AtPAT1), and dwarf and low tillering (DLT) (Sun et al., 2011). The SCR and SHR subfamilies were validated to act as positive regulators to promote the formation of root radial morphology and root monolayer cells in *Arabidopsis* (Koizumi et al., 2012; Rich et al., 2017). SCR is mainly expressed in root epidermal cells regulating the division of daughter cells differentiation in root, whereas SHR not only affects the polar differentiation of root endothelial cells but also activates the SCR promoter in specific tissues to regulate its function (Slewinski et al., 2012). In addition, SHR proteins are expressed explicitly in the mid-column sheath cells. They can transfer from external monolayers to endothelial cells to form a dimer with SCR, which jointly activates the transcription of the *SCR* gene and other downstream target genes. Therefore, it can be inferred that SCR and SHR are transcriptionally dependent on each other, and the resulting SCR-SHR complex positively modulates the transcription of *SCR* (Koizumi et al., 2012; Slewinski et al., 2012). Yeast extract (YE), also known as yeast flavoring, plays a vital role in promoting the accumulation of secondary metabolites in plants. In *S. miltiorrhiza*, after treating hairy root culture with YE, the content of tanshinone was significantly increased (Chen et al., 2001). However, the transcriptional mechanism involved in YE-triggered tanshinone biosynthesis remains obscure.

In this study, we isolated and identified a new GRAS TF named *SmSCR1*. Our studies uncover the regulatory role of *SmSCR1* gene in response to YE-induced regulation of tanshinone biosynthesis and hairy root growth and development. Our data discover a novel TF family, GRAS, as a candidate for improving the biosynthesis potential of tanshinone through metabolic engineering strategies.

**MATERIALS AND METHODS**

**Plant Materials and Growth Conditions**

*Salvia miltiorrhiza* plants were grown in solid Murashige and Skoog (MS) medium at 25°C, in 60% humidity, and in a light rhythm of 16-h light and 8-h dark (Shi et al., 2021a,b). *S. miltiorrhiza* hairy roots were subcultured in 250 ml Erlmeyser flasks with 100 ml 1/2 MS liquid medium and placed in a shaker at 110 rpm min⁻¹, 25°C in the dark. The hairy roots were collected after 60 days for gene expression analysis (Shi et al., 2016a, 2020). *Nicotiana benthamiana* was directly sown in soil and cultivated in pots for further experimental needs (Huang et al., 2019). RNA was extracted from different tissues of annual plants of *S. miltiorrhiza* and stored at −80°C using liquid nitrogen flash freezing.

**Gene Isolation and Sequence Analysis**

All the GRAS families in *S. miltiorrhiza* were identified against local datasets. *SmSCR1*, which belongs to the GRAS family and intensively responds to YE induction, was cloned using a homology-based cloning method, as described previously (Zhou et al., 2016, 2017). The cloned *SmSCR1* gene was analyzed using the BlastP tool in the NCBI database. The reported GRAS members, i.e., *SmGRAS1* (accession number: KY435886) and *SmGRAS2* (KY435887) in *S. miltiorrhiza*, together with *O. sativa* OsGRAS32 (Os06g0127800), *O. sativa* OsSCR1 (XP_015615402.1), *O. sativa* OsSCL2 (XP_015620600.1), *A. thaliana* AtSCR (Q9M384.1), and *A. thaliana* AtSCL23 (NM_123557), which are highly homologous to *SmSCR1* (OM032820), were subjected to amino acid sequence alignment using Vector NTI software (Invitrogen, Carlsbad, CA, United States). The full-length amino acid sequences of the GRAS were aligned using the neighbor-joining (NJ) method in the Clustal X computer, and the MEGA 6.0 program was used to generate a phylogenetic tree. Using Clustal X computing tools, the amino acid sequences of candidate GRAS family members were aligned based on the NJ method, and MEGA 6.0 software was used to generate phylogenetic trees. Subfamily members of SCR and SCL from *Arabidopsis* and rice were compared with *SmSCR1* using Vector NTI 10.0 software.

**Subcellular Localization Analysis of SmSCR1**

To verify the subcellular localization profile of *SmSCR1*, pHB-*SmSCR1*-YFP, and pHB-YFP recombinant plasmids were

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**Supplementary Figure 1**
introduced into Agrobacterium rhizogenes strain EHA105 for transient transformation, respectively. pHB-YFP was used as the positive control. After 48 h of infection by Agrobacterium tumefaciens, YFP signals from infected N. benthamiana leaves were visualized using a high-resolution microscope observation system. The nuclei of epidermal cells of infected N. benthamiana leaves were stained with 4′, 6-diamidino-2-phenylindole dihydrochloride (DAPI) solution for 3 h before observation (Zhou et al., 2020).

Quantitative Real-Time PCR
According to the procedure reported before (Shi et al., 2016b), the quantitative real-time PCR (qRT-PCR) analysis was conducted. Different tissues (e.g., taproot, lateral root, fibrous root, phloem, xylem, stem, petiole, young leaves, and mature leaves) were taken from an adult plant, as well as hairy roots with the treatment of YE, and then frozen in liquid nitrogen for RNA isolation (Zhang et al., 2011). Actin gene of S. miltiorrhiza was used as the internal reference. The sequences of all primer pairs used in qRT-PCR studies are summarized in Supplementary Table 1. The gene expression was quantified by the comparative Ct method (Hao et al., 2015).

Transformation of SmSCR1 in Salvia miltiorrhiza
The open reading frame (ORF) of SmSCR1 was introduced into the double restriction insertion site of BamHI and SpeI of the pH8 vector under the CaMV 35S promoter and NOS terminator (Supplementary Figure 2A). The DNA sequence encoding the SRDX structural domain (LDLDLELRLGFA) was inserted into the ORF of SmSCR1 just before the stop codon (TAA) using the method described previously (Hiratsu et al., 2003; Deng et al., 2020a,b). The SmSCR1-SRDX was then subcloned into the pHB to create the pH8-SmSCR1-SRDX suppressive expression construct (Supplementary Figure 2B). To obtain the transgenic hairy roots, all plasmids were transformed into...
FIGURE 2 | Multiple alignments of SmSCR1 with related GRAS proteins from other plants. SmGRAS1 (accession number: KY435886), SmGRAS2 (KY435887), OsGRAS32 (Os06g0127800), OsSCR1 (XP_015615402.1), OsSCR2 (XP_015620600.1), AtSCR (Q9M384.1), AtSCL23 (NM_123557), and SmSCR1 (OM032820). Black line indicates the conserved amino acid sites. ILR I, leucine-rich region I; LR II, leucine-rich region II; VHIID, the conserved region consisting of valine, histidine, isoleucine, and D-aspartic acid; PFYRE, the conserved amino acid sites consisting of proline, phenylalanine, tyrosine, arginine, and glutamic acid; SAW, the conserved region consisting of serine, alanine, and tryptophan.
A. tumefaciens C58C1 and then transinfected into S. miltiorrhiza (Zhou et al., 2016).

**Determination of Tanshinones by High-Performance Liquid Chromatography**

After 60 days of continuous culture in 1/2 MS liquid medium, each hairy root line was harvested and dried in a freeze dryer for 24 h, and then extracted with 16 ml methanol/dichloromethane (3:1, v/v). Notably, 100 mg of dried powder of S. miltiorrhiza, sonicated for 1 h, then soaked overnight in the dark and centrifuged for 10 min at 12,000 × g. After centrifugation, the supernatant was poured into a distillation flask. After centrifugation, the supernatant was poured into a distillation flask and spun dry at 50°C. We then added 2 ml of methanol to dissolve the material in the distillation flask. After further centrifugation at 6,500 × g for 10 min, the samples were filtered separately through 0.22 μM organic membranes (Jinten, China) using a 1 ml sterile syringe (Huang et al., 2019). All metabolites were quantified by high-performance liquid chromatography (HPLC), as described previously (Agilent Technologies, Santa Clara, CA, United States) (Zhou et al., 2016; Sun et al., 2019; Yang et al., 2021).

**Dual-Luciferase Assay**

The dual-luciferase (Dual-LUC) assays were carried out as previously reported (Deng et al., 2019). The recombinants of pHBDYFP and pHBSmSCR1-YFP were introduced into A. tumefaciens GV3301, respectively. Gene promoters from the tanshinone biosynthetic pathway were cloned and inserted into
pGREEN0800 vector, respectively, which were co-transformed into the GV3301 strain with pSoup19. The test was carried out as previously reported (Cao et al., 2018). The pHB-YFP construct was employed as the negative control, and the Renilla gene was used as the internal control.

**Yeast One-Hybrid Assay**

Similar to a previous study (Deng et al., 2019), the ORF of SmScr1 was inserted into the pB42AD vector. Three fragments of SmCps1 promoter sequence (i.e., from −1 to −700, −701 to −1,400, and −1,401 to −2,100 bp, relative to translation start site, respectively) were inserted into a pLacZ plasmid. The resulting recombinants were co-transformed into yeast cell EGY48a. After cultivation on SD/-Ura/-Leu medium for 48 h, the positive binding activity was examined on SD/-Ura/-Leu medium with X-gal for 24 h. PB42AD and pLacZ empty carriers were set as negative controls.

**Morphological Observations of SmScr1**

To investigate the effect of the SmScr1 gene on plant growth and development, we examined different phenotypes of the transgenic hairy roots on 1/2 MS medium. In addition, three different transgenic hairy roots of SmScr1 with good growth and similar developmental time were selected for developmental microstructure observation, as reported previously (Wu et al., 2020). The pith (Pi) and posterior xylem (Mx) of the hairy roots were observed under a light microscope (200 ×).

**Statistical Analysis**

Each data represent the average of three independent experimental calculations, and the data are expressed as the mean ± SD. SPSS 16.0 software was used to perform a single-sample t-test and one-way ANOVA, and p-values < 0.05 were considered statistically significant.

**RESULTS**

**Isolation and Sequence Analysis of SmScr1 Gene**

First, we identified a GRAS TF (unigene number: SmiContig9465) and SmScr1 (NCBI accession number:...
OM032820), responding to YE-induction (fold change > 2) from the *S. miltiorrhiza* transcriptome (Zhou et al., 2017). *SmSCR1* length is 1,944 bp long and encodes 647 amino acids, with a molecular weight size of 70,515 kDa coupled with a theoretical isoelectric point of 5.78. GRAS family members from *Arabidopsis* and rice were analyzed with *SmSCR1* to construct a phylogenetic tree. As shown in Figure 1, *SmSCR1* has the highest relationship with AtSCR in *Arabidopsis* and OsSCR2 in rice, implying that *SmSCR1* belongs to the SCR subgroup of the GRAS family. *SmSCR1* consists of several conserved motifs, including VHIID, PFYRE, SAW motifs, and the leucine-rich regions (LRs), i.e., LR I and LR II (Figure 2).

**Expression Analysis of SmSCR1**

Transient expression of *SmSCR1* in epidermal cells from 45-day-old *N. benthamiana* leaves exhibited that *SmSCR1* located in the nucleus (Figure 3A). The gene expression of *SmSCR1* was characterized from nine different tissues of *S. miltiorrhiza* by qRT-PCR. *SmSCR1* was expressed in all tissues, namely, taproot, lateral root, fibrous root, phloem, xylem, stem, petiole, young leaves, and mature leaves, with the highest expression in xylem and petiole, and the lowest expression in taproot (Figure 3B). The expression of *SmSCR1* after YE-induction was examined, reaching the highest levels for 2 h after post-induction with 13.6-fold compared to the control (Figure 3C).

**SmSCR1 Activates the Transcription of Copalyl Diphosphate Synthase 1 Gene**

The role of *SmSCR1* in regulating tanshinone biosynthetic gene expressions was examined. The Dual-LUC assay showed that *SmSCR1* uniquely activates the transcription of the *SmCPS1* promoter, leading to a 2.19-fold increase compared with the control, but not to other promoters (e.g., *HMGR1* promoter) (Figure 4). YH assay showed that self-activating of pB42AD vector binding to the promoter of the CPS1 gene is visible (Supplementary Figure 3). Those results indicated that *SmSCR1* activated the transcription of *SmCPS1* to increase the production of tanshinone in *S. miltiorrhiza*.

**SmSCR1 Promotes Tanshinone Biosynthesis in the Transgenic Hairy Root of Salvia miltiorrhiza**

The OE and SRDX suppressive hairy root lines were first examined by genomic PCR detection (Supplementary Figure 4). Then, the transcript profiles of *SmSCR1* in transgenic hairy roots were quantified by qRT-PCR. Three independent OE lines (i.e., OE-5, OE-8, and OE-11) and three SRDX lines (i.e., SRDX-2, SRDX-4, and SRDX-6) of *SmSCR1* with the highest expression levels in transgenic lines were subsequently chosen for further analysis (Supplementary Figure 5). After 60-day-old subculture, the hairy root lines were collected independently and were subjected to examine the content of tanshinones by HPLC (Figure 5A). The concentrations of DT, TA-1, CT, and TTA were significantly increased compared to the control in the OE lines, with the highest total tanshinone (TTA) content in *SmSCR1*-OE-11, reaching 1.49-fold than that of the control. In contrast, in the SRDX repression lines, DT, TA-1, CT, and TTA were significantly decreased compared to the control, with the lowest TTA content of 2.76 mg/g DW. The above results suggest that the *SmSCR1* TF can promote the tanshinone accumulation in *S. miltiorrhiza*.

**SmSCR1 Upregulates Copalyl Diphosphate Synthase 1 and HMGR1 in Transgenic Hairy Roots of Salvia miltiorrhiza**

Copalyl diphosphate synthase 1 and *HMGR1* were significantly upregulated in OE lines (e.g., OE-5, OE-8, and OE-11), with the *SmSCR1*-OE-11 line, showing the highest expression. In contrast, the two genes were drastically downregulated in the three SRDX lines (i.e., 2, 4, and 6), with the *SmSCR1*-SRDX-6 line exhibiting the lowest decrease compared to the control (Figure 5B). These findings showed that *SmSCR1* stimulated *CPS1* and *HMGR1* together, resulting in increased production in transgenic hairy roots of *S. miltiorrhiza*.

**SmSCR1 Promotes the Growth of Transgenic Hairy Roots of Salvia miltiorrhiza**

The biomass of the transgenic hairy root lines significantly increased in the *SmSCR1* OE lines and significantly reduced in the SRDX lines compared to the control (Figure 6A). Moreover, the *SmSCR1* overexpression hairy root lines had a better growth status than the control, with more branches in the *SmSCR1* OE lines and fewer in the SRDX suppressive hair roots (Figure 6B). Roots are formed by a series of asymmetrical divisions of early differentiated cells to produce the inner and outer cortex of the root, which then develops into xylem and phloem. The development of transgenic hairy roots by paraffin sectioning demonstrated that the *SmSCR1* gene promotes the formation of hairy root radiation morphology of the Mx cell and Pi cell differentiation in *SmSCR1*-OE hairy roots when compared with the control and the SRDX suppressive hairy roots (Figure 6). Those results suggest that *SmSCR1* can modulate the growth and development of hairy roots in *S. miltiorrhiza*.

**DISCUSSION**

The GRAS TF family is an important TF class involved in regulating plant growth, development, environmental stress response, and growth signal transduction (Sun et al., 2011). With the development of genome sequencing technology, a few GRAS family members have been identified in plants, such as plum (*Armeniaca mume*), lotus (*Nelumbo nucifera*), and pine blue (*Isatis indigotica*) (Kamiya et al., 2003; Hou et al., 2010). However, only a few reports systematically characterized GRAS TFs in medicinal plants. We identified a novel YE-responsive GRAS TF gene, *SmSCR1* gene, encoding 647 amino acids with several key GRAS conserved structural domains (e.g., VHIID, LR I, LR II, PFYRE, and SAW motifs) (Koizumi et al., 2011; Zhou et al., 2017). In a previous study, SCR and SHR families
were thought to co-regulate the plant growth, development, and radial structures in roots (Koizumi et al., 2011, 2012; Fan et al., 2017). We also identified that SmSCR1 was very sensitive to YE-induction (Figure 3C), regulating tanshinone accumulation in *S. miltiorrhiza* (Zhou et al., 2017, 2021a).

We revealed that SmSCR1 upregulates tanshinone biosynthesis in SmSCR1 suppressive hairy root lines (Figure 5A). Recently, two GRAS families, including SmGRAS1 (GenBank accession number: KY435886) and SmGRAS2 (GenBank accession number: KY435887), clustered into SHR and PAT1 subgroups, respectively (Figure 1), were characterized in *S. miltiorrhiza*. SmGRAS1 and SmGRAS2 were verified by transgenic validation to upregulate tanshinone accumulation in hairy roots. Therefore, many GRAS families might play a more prominent role in regulating tanshinone accumulation in *S. miltiorrhiza* and require further investigation.

Both CPS1 and HMGR1 positively correlated to the expression of SmSCR1 in the transgenic lines, while SmSCR1 can only bind to the CPS1 promoter to activate its expression as examined. Due to the limited data about the specific *cis*-element in promoter binding by SCR1 in other plants (Li et al., 2019), further examining the direct binding of SmSCR1 to the specific...
promoter elements by EMSA assay \textit{in vitro} would be unfeasible. In this study, overexpressing \textit{SmSCR1} upregulated not only the expression of its target gene \textit{CPS1} but also the \textit{HMGR1} gene in the tanshinone biosynthetic pathway. Overexpressing \textit{SmERF1L1} promotes the expression of five genes in the tanshinone biosynthetic pathway, but only \textit{DXR} is a target (Huang et al., 2019). In \textit{SmERF128} transgenic hairy roots, the expression of six genes in the tanshinone biosynthetic pathway was activated, and only three of the six genes were validated to be targets of \textit{SmERF128} (Zhang et al., 2019). Thus, it can be inferred that the ectopic expression of certain TF genes upregulates the expression of not only their target genes but also other genes in the same biosynthetic pathway.

The root formation is a series of asymmetric divisions of stem cells to generate the inner and outer cortex of the root and then differentiate into different forms of tissue. In \textit{A. thaliana}, SCR is mainly expressed in root epidermal cells. As a positive regulator, SCR regulates root radiation morphology and plays a vital role in root monolayer cell formation (Koizumi et al., 2012; Rich et al., 2017). When SCR function is lost and mutated, the normal development of the cell layer will be affected. In addition, SCR can also regulate the division of progenitor cells and promote the growth of biomass (Koizumi et al., 2011; Slewinski et al., 2012). In rice, \textit{OsSCR} was thought to affect the development of plant leaf guard cells (Hou et al., 2010). In the three \textit{SmSCR1} OE lines, we found that \textit{SmSCR1} significantly promoted the biomass increase of the three transgenic hairy root lines, whereas a visible decrease was observed in \textit{SmSCR1} SRDX lines.

Moreover, \textit{SmSCR1} can regulate the development of \textit{Pi} and the \textit{Mx}. In \textit{SmSCR1} OE lines, the cell division rate of progeny

![FIGURE 7](image1.png)

**FIGURE 7** | Paraffin sectioning analysis of the transgenic hairy roots. The control means the transgenic hairy roots with the transformation of pHB-X-YFP blank vector. pHB-SmSCR1 represents the overexpression lines, and pHB-SmSCR1-SRDX represents the suppression lines. \textit{Pi} represents the pith. \textit{Mx} represents the posterior xylem.

![FIGURE 8](image2.png)

**FIGURE 8** | A proposed model for the role of \textit{SmSCR1} in modulating tanshinone biosynthesis and root growth in \textit{Salvia miltiorrhiza}.
and the root radial structure growth rate were significantly higher than that of control. In contrast, the three SRDX suppressive hairy root lines have an opposite phenotype (Figure 7). The function of SmSCR1 to regulate the growth and development of hairy roots in S. miltiorrhiza is similar to the model plants, e.g., Arabidopsis and rice.

Based on the present findings, a deductive model of SmSCR1 as an activator in response to YE-induction promotes tanshinone accumulation, Pi cell division, radial structure formation, and biomass increase of hairy roots in S. miltiorrhiza was outlined (Figure 8).

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author.

AUTHOR CONTRIBUTIONS

GK and WZ conceived and designed the project. SW, YS, YL, XG, CC, and TL prepared materials and performed the experiments. WZ, SW, YS, IM, YL, and CW performed the bioinformatics analysis and analyzed the data. GK, WZ, SW, and YS wrote the manuscript. IM, WZ, and GK revised the manuscript. All authors read and approved the final version of the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2022.860033/full#supplementary-material

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