Biosyntheses characterization of alkaloids and flavonoids in *Sophora flavescens* by combining metabolome and transcriptome

Guangfei Wei¹, Yongzhong Chen¹, Xiaotong Guo², Jianhe Wei³, Linlin Dong¹,* & Shilin Chen¹,*

*Sophora flavescens* are widely used for their pharmacological effects. As its main pharmacological components, alkaloids and flavonoids are distributed in the root tissues wherein molecular mechanisms remain elusive. In this study, metabolite profiles are analyzed using metabolomes to obtain biomarkers detected in different root tissues. These biomarkers include alkaloids, phenylpropanoids, and flavonoids. The high-performance liquid chromatography analysis results indicate the differences in principal component contents. Oxymatrine, sophoridine, and matrine contents are the highest in the phloem, whereas trifolirhizin, maackiain, and kushenol I contents are the highest in the xylem. The transcript expression profiles also show tissue specificity in the roots. A total of 52 and 39 transcripts involved in alkaloid and flavonoid syntheses are found, respectively. Among them, the expression levels of LYS*A1, LYS*A2, A02, A06, PMT1, PMT17, PMT34, and PMT35 transcripts are highly and positively correlated with alkaloids contents. The expression levels of 4CL1, 4CL3, 4CL12, CHI5, CHI7, and CHI9 transcripts are markedly and positively correlated with flavonoids contents. Moreover, the quantitative profiles of alkaloids and flavonoids are provided, and the pivotal genes regulating their distribution in *S. flavescens* are determined. These results contribute to the existing data for the genetic improvement and target breeding of *S. flavescens*.

Abbreviations

UPLC-MS Ultrahigh-performance liquid chromatography quadrupole time of flight-mass spectrometry
HPLC-UV High-performance liquid chromatography-ultraviolet detection
PCA Principal component analysis
ANOVA One-way analysis of variance
VIP Variable importance in projection
FDR False discovery rate
KEGG Kyoto encyclopedia of genes and genomes
FPKM Fragments per kilobase of exon per million fragments
DEGs Differential expression genes
NCBI Nr NCBI non-redundant protein
WGCNA Weighted gene co-expression network analysis
UGTs UDP-glycosyltransferases
P450 P450-monooxygenase
C4H Cinnamate 4-hydroxylase
4CL 4-Coumaric acid coenzyme A ligase
CHS Chalcone synthase
CHI Chalcone isomerase
PAL Phenylalanine/tyrosine ammonia-lyase

¹Key Laboratory of Beijing for Identification and Safety Evaluation of Chinese Medicine, Institute of Chinese Materia Medica, China Academy of Chinese Medical Sciences, Beijing 100700, China. ²College of Agriculture, Ludong University, Yantai 264025, China. ³Hainan Provincial Key Laboratory of Resources Conservation and Development of Southern Medicine, Hainan Branch of the Institute of Medicinal Plant Development, Chinese Academy of Medical Sciences and Peking Union Medical College, Haikou 570311, China. ⁴Present address: No.16 Nanxiaojie, Dongzhimennei Ave., Beijing 100700, China. *email: lldong@icmm.ac.cn; slchen@icmm.ac.cn
The alkaloids contents in the root tissues is as follows: phloem > xylem > pith > cork layer. The biosynthesis of active ingredients in medicinal plants is often related to the synergistic expression and regulation of key enzyme genes. Transcriptomes have been used to study the transcription and expression levels of genes in medicinal plants. For example, a total of 749 ginsenoside biosynthetic enzyme genes, together with 12 good pleiotropic drug resistance genes related to ginsenoside transport, were identified from the adventitious roots of *Panax ginseng* and *Panax quinquefolius*, respectively. More than 200 compounds have been isolated and identified from *Sophora* Radix, including alkaloids, flavonoids, terpenoids, and other compounds. Their composition and content vary between the organs of *Sophora* Radix, such as the roots, stems, leaves, flowers, and seeds. However, the reports on the distribution of alkaloids and flavonoids in the root tissues of *Sophora flavescens* are rare. Therefore, the difference in the profiles of alkaloids and flavonoids components in the root tissues of *S. flavescens* should be established for the targeted breeding of this species.

In negative ion mode, 11,101 components were detected, among which 297 components were identified (Supplemental file 2: Dataset S2). The PCA results also revealed a clear separation between the three root tissues of *S. flavescens* in positive ion mode (Fig. 1a). A total of 387 potential biomarkers were detected in negative ion mode through ANOVA analysis (FDR ≤ 0.05; Fig. 1e and Supplemental file 2: Dataset S2). The PCA results also revealed a clear separation between the three root tissues of *S. flavescens* in positive ion mode (Fig. 1a). A total of 387 potential biomarkers were detected in negative ion mode through ANOVA analysis (FDR ≤ 0.05; Fig. 1e and Supplemental file 2: Dataset S2). The PCA results also revealed a clear separation between the three root tissues of *S. flavescens* in positive ion mode (Fig. 1a). A total of 387 potential biomarkers were detected in negative ion mode through ANOVA analysis (FDR ≤ 0.05; Fig. 1e and Supplemental file 2: Dataset S2).
M301T3251 (Moracin M), M269T239 (Aloe-emodin), M313T285 (Velutin), and M251T170 (Ser-Phe) were the most abundantly present in the xylem tissue (VIP > 1; Fig. 1f and Supplemental file 2: Dataset S2). These data showed that metabolites existed differences in the root tissues of *S. flavescens*.

**Distribution of alkaloids and flavonoids in the root tissues of *S. flavescens***. The contents of the three alkaloids (oxymatrine, sophoridine, and matrine) and five flavonoids (trifolirhizin, maackiain, kushenol I, 

---

**Figure 1.** Metabolomic analysis of the components in the root tissues of *S. flavescens*. (a) PCA score plots of positive ion mode. (b) One-way ANOVA of the positive ion mode. (c) Variable importance in the projection of positive ion mode. (d) PCA score plots of negative ion mode. (e) One-way ANOVA of negative ion mode. (f) Variable importance in the projection of negative ion mode.
kurarinone, and sophoraflavanone G) were detected through HPLC (Fig. 2). The contents of the three alkaloids were the highest in the phloem (23.93, 1.88, and 1.83 mg/g, respectively), followed by the xylem (17.73, 0.56, and 1.40 mg/g, respectively) and periderm (11.97, 1.04, and 0.21, respectively; Fig. 2a). The contents of trifolirhizin, maackiain, and kushenol I were the highest in the xylem (0.21, 0.49, and 1.31 mg/g, respectively). Finally, kurarinone and sophoraflavanone G contents were the highest in the periderm (0.01 and 0.01 mg/g, respectively; Fig. 2b). These results showed that alkaloids and flavonoids existed distribution differences in the root tissues of S. flavescens.

Transcriptome profiles in the root tissues of S. flavescens. Illumina Hiseq paired-end sequencing technology was used to analyze the transcriptome in the root tissues of S. flavescens (Table 1). The average raw reads was 7.61 G, and the average clean reads was 6.81 G after filtered by fqtrim software. The percentages of Q20, Q30, and GC were 98.35%, 94.89%, and 45.83%, respectively. All 296,523,264 high-quality 150 bp clean reads were used for de novo assembly, and in total 35,012 contigs > 500 bp in length were obtained (Table 2). Then, a total of 58,327 genes were assembled, with an N50 contig size of 1,237 bp. Annotation was also performed on the basis of the sequence similarity searches with a cutoff E-value of 10^{-5} against public databases, including the GO, Kyoto Encyclopedia of Genes and Genomes (KEGG), Pfam, SwissProt, eggNOG, and Nr databases, to investigate the function of assembled unigenes (Table 3). A total of 24,549 (66.63%), 19,056 (51.72%), 21,697

![Figure 2. Contents of alkaloids and flavonoids in the root tissues of S. flavescens. (a) Contents of three alkaloids. (b) Contents of five flavonoids. Pe, Ph, and Xy represent the periderm, phloem, and xylem, respectively.](https://example.com/figure2.png)

| Samples  | Raw reads | Raw bases | Clean reads | Clean bases | Q20%  | Q30%  | GC%  |
|----------|-----------|-----------|-------------|-------------|-------|-------|------|
| Periderm 1 | 56,056,078 | 8.41G | 54,669,246 | 7.55G | 98.69 | 95.55 | 45.28 |
| Periderm 2 | 51,184,822 | 7.68G | 49,657,472 | 6.82G | 98.17 | 94.45 | 47.27 |
| Periderm 3 | 45,266,188 | 6.79G | 43,970,534 | 6.07G | 98.37 | 95.16 | 46.50 |
| Phloem 1   | 49,555,614 | 7.43G | 48,015,280 | 6.60G | 98.23 | 94.65 | 46.44 |
| Phloem 2   | 63,603,050 | 9.54G | 61,977,910 | 8.54G | 98.31 | 94.78 | 45.33 |
| Phloem 3   | 50,063,620 | 7.51G | 48,836,916 | 6.73G | 98.37 | 94.93 | 45.47 |
| Xylem 1    | 45,639,898 | 6.85G | 44,549,806 | 6.14G | 98.35 | 94.89 | 45.67 |
| Xylem 2    | 47,552,830 | 7.13G | 46,387,724 | 6.39G | 98.30 | 94.75 | 45.15 |
| Xylem 3    | 47,837,988 | 7.18G | 46,720,006 | 6.44G | 98.34 | 94.87 | 45.34 |
| Average    | 50,751,121 | 7.61G | 49,420,544 | 6.81G | 98.35 | 94.89 | 45.83 |

Table 1. Statistics of transcriptome data for S. flavescens.

| Item                | No. of sequences |
|---------------------|-----------------|
| High-quality reads  | 296,523,264     |
| No. of contig > 500 bp | 35,012          |
| Total unigenes      | 58,327          |
| N50 contig size (bp)| 1,237           |
| Total assemble bases (bp) | 50,280,338     |

Table 2. Summary of the transcriptome data and the assembly results of Sophora flavescens.
(58.89%), 20,069 (54.47%), 27,094 (73.54%), and 27,647 (75.04%) unigenes had significant matches with the GO, KEGG, Pfam, SwissProt, eggNOG, and NR databases, respectively.

**Gene expression profiles in the root tissues of S. flavescens.** The PCA and Venn profiles were carried out to investigate the transcription distinction among the main root tissues in S. flavescens on the basis of the Fragments Per Kilobase of exon per Million fragments (FPKM) value (Fig. 3). The PCA results showed that the three root tissues existed slightly difference (Fig. 3a). The Venn results showed that 5129 unigenes were shared and expressed among the three tissues. A total of 3454, 3666, and 4117 unigenes were explicitly defined in the periderm, phloem, and xylem, respectively (Fig. 3b). With the comparison between the periderm and the phloem and xylem, 82 (Up: Down, 63: 19) and 654 (Up: Down, 454: 200) DEGs were found, respectively (Supplemental file 1: Figure S2). With the comparison between the phloem and the xylem, 186 (Up: Down, 45: 141) DEGs were found. These results showed that the unigenes in the root tissues of S. flavescens had different expression levels.

**Co-expression analysis of the transcripts and active components in S. flavescens.** In S. flavescens, all transcripts were grouped into 23 unique modules, and two modules, namely, MEmagenta and MElightyellow, were positively correlated with active components (\( R < 0.05; \) Fig. 4a). MEmagenta was significantly and positively correlated with the contents of trifolirhizin (\( R = 0.73 \)) and maackiain (\( R = 0.71 \); Fig. 4b). MElightyellow was positively correlated with the contents of kurarinone (\( R = 0.88 \)).

**Analysis of the transcripts involved in the alkaloid and flavonoid biosyntheses in the root tissues of S. flavescens.** Key transcripts and enzymes resulted in various regulatory controllers in the alkaloid and flavonoid biosyntheses. The expression of most transcripts significantly differed (Fig. 5 and Supplemental file 2: Dataset S3, S4, and S5). In the alkaloid upstream biosynthesis pathway, 52 transcripts were selected to analyze expression profiles. Moreover, five (9.62%), 16 (30.77%), and 31 (59.62%) transcripts were expressed at the highest levels in the periderm, phloem, and xylem, respectively (Fig. 5a and Supplemental file 2: Dataset S3). One DMR transcript (DMR1), one AO transcript (AO2), and three PMT transcripts (PMT10, PMT11, and PMT19) showed the highest expression levels in the periderm. One LYSA (LYSA3), two DMR (DMR2 and DMR3), seven AO (AO1, AO3, AO4, AO7, AO8, AO9, and AO10), and 23 PMT (PMT2, PMT3, PMT4, PMT6, PMT8, PMT12, PMT13, PMT14, PMT15, PMT16, PMT18, PMT19, PMT20, PMT23, PMT24, PMT25, PMT26, PMT27, PMT28, PMT29, PMT31, PMT33, and PMT36) had the highest expression levels in the xylem. A total of 137 CYP transcripts participated in alkaloid synthesis were identified. Among them, 14 (10.22%), 52 (37.96%), and 71 (51.82%) transcripts were expressed at the highest levels in the periderm, phloem, and xylem,

| Total | Nr   | Nt   | Swiss-Prot | KEGG  | COG   | GO    | Overall |
|-------|------|------|------------|-------|-------|-------|---------|
| 84,408| 68,720| 49,929| 52,118     | 52,176| 28,431| 42,535| 69,951  |

100% 81.88% 61.29% 63.65% 63.54% 36.02% 52.80% 83.10%

**Table 3.** The annotation of all unigenes in *Sophora flavescens.*

**Figure 3.** (a) Principal component analysis (PCA) and (b) Venn profiles in the roots of *S. flavescens.*
Figure 4. Co-expression profiles of all transcripts and active components in the root tissues of *S. flavescens*. (a) Hierarchical cluster tree showing co-expression modules in *S. flavescens*. (b) Module-components association in *S. flavescens*. 
respectively (Supplemental file 1: Figure S3 and Supplemental file 2: Dataset S4). These results showed that the transcripts related to alkaloid synthesis were highly expressed in the xylem.

In the flavonoid biosynthesis pathway, 39 transcripts were selected to analyze expression profiles, wherein 5 (12.83%), 8 (20.5%), and 26 (66.67%) transcripts yielded the highest expression levels in the periderm, phloem, and xylem, respectively (Fig. 5b and Supplemental file 2: Dataset S5). Three 4CL transcripts (4CL1, 4CL7, and 4CL12) and one 2′OH transcript (2′OH2) exhibited the highest expression levels in the periderm. Three C4H (C4H1, C4H2, and C4H3), 11 CL (CL2, CL3, CL4, CL5, CL6, CL8, CL9, CL10, CL11, CL14, and CL15), 5CHS (CHS1, CHS2, CHS3, CHS4, CHS5), seven CHI (CHI1, CHI2, CHI3, CHI4, CHI6, and CHI8), and one 2′OH (2′OH3) yielded the highest expression levels in the xylem. These results showed that the transcripts related to flavonoid synthesis were highly expressed in the xylem.

Correlation analysis of the active component contents and transcript expressions in *S. flavescens*. The alkaloid synthetic pathway showed high and positive associations between component contents and transcripts: 16 transcripts with oxymatrine contents, two transcripts with sophoridine contents, and 24 transcripts for matrine contents \((R > 0.8, P < 0.05, \text{Fig. 6a})\). The expression levels of two LYS (LYS1 and LYS2), two AO (AO2 and AO6), and 12 PMT (PMT1, PMT5, PMT7, PMT9, PMT17, PMT21, PMT22, PMT25, PMT31, PMT34, and PMT35) transcripts were markedly and positively correlated with oxymatrine contents. The expression levels of two PMT (PMT7 and PMT30) transcripts were highly correlated with sophoridine contents. The expression levels of two LYSA (LYSA1 and LYSA2), three AO (AO2, AO6, and AO9), and 19 PMT (PMT1, PMT2, PMT5, PMT6, PMT7, PMT9, PMT12, PMT17, PMT21, PMT22, PMT25, PMT26, PMT28, PMT31, PMT32, PMT33, PMT34, PMT35, and PMT36) transcripts were markedly and positively correlated with matrine contents.

In the flavonoid synthesis pathway, a total of 3, 3, 4, and 4 transcripts were highly and positively correlated with the contents of trifolirhizin, maackiain, kushenol I, kurarinone, sophorafavanone G, respectively \((R > 0.8, P < 0.05, \text{Fig. 6b})\). The expression levels of two 4CL (4CL1 and 4CL12) and one 2′OH (2′OH2) transcripts were markedly and positively correlated with trifolirhizin and maackiain contents. The expression levels of two 4CL (4CL1 and 4CL13) and one 2′OH (2′OH1) transcripts were highly correlated with kuslenol I contents. The expression levels of CHI5 and CHRI1 transcripts were highly correlated with kurarinone and sophorafavanone G contents.

Discussion

In this study, the distribution of alkaloids and flavonoids demonstrated tissue specificity in *S. flavescens* roots. Transcript expression profiles also existed tissue specificity in the roots. The weighted gene co-expression network analysis (WGCNA) results confirmed that the pivotal transcripts regulated the distribution of alkaloids and flavonoids in the root tissues. This study will provide useful information for investigating the genetic and biochemical mechanisms of alkaloid and flavonoid syntheses.
Metabolite profiles revealed that the chemical components showed tissue specificity in the root tissues of *S. flavescens*. A total of 387 and 257 biomarkers were detected in positive and negative ion modes, respectively. The biomarkers detected in this study included many components of alkaloids and flavonoids. Alkaloids and flavonoids are the main chemical components of *S. flavescens*. They possess significant pharmacological effects, such as anti-tumor and anti-virus activities (matrine)\(^1\), hypoglycemic and hypolipidemic effects (oxymatrine)\(^1\), human colorectal cancer preventions (sophoridine)\(^2\), anti-proliferation (trifolirhizin)\(^3\), and inflammasome-activating effect (maackiain)\(^4\). Thus, the contents of alkaloids and flavonoids in the roots of *S. flavescens* were further quantitatively analyzed. Quantitative analysis results revealed that the contents of the total alkaloids and three alkaloids (oxymatrine, sophoridine, matrine) were higher in the phloem than those in the periderm and

---

**Figure 6.** Pearson correlation bubble chart of the transcript expression and chemical component contents in *S. flavescens*: (a) alkaloids and (b) flavonoids. The size of the circle represents the correlation coefficient. The color red represents a positive correlation, whereas the color green represents a negative correlation.
xylem. This finding was consistent with the reports of previous work. The contents of trifolirhizin, maackiain, and kushenol I were the highest in the xylem, and the contents of kurarinone and sophorallavananone G were the highest in the periderm. Four phenolic acid compounds (benzoic acid, caffeic acid, ferulic acid, and chlorogenic acid) and four flavonol compounds (kaempferol, catechin hydrate, epicatechin, and rutin) were higher in the aerial parts than the roots. This uneven accumulation pattern of secondary metabolites may affect the rational use of medicinal plants. Understanding the molecular biological mechanism of the active components is of great significance.

The transcript expression in *S. flavescens* was tissue-specific. A total of 52 upstream transcripts and 137 downstream CYP transcripts involved in alkaloid synthesis were identified (FPKM ≥ 5), among which 59.62% and 51.82% were expressed at the highest levels in the xylem. In a previous study, the preferential expression of the gene for putative lysine/ornithine decarboxylase committed in the initial step of matrine biosynthesis was the highest in the leaf and stem. The above finding evidentially indicated that the different expressions of these genes resulted in the uneven distribution of alkaloids. In the flavonoid compound biosynthesis pathway, 26 (66.67%) transcripts were highly expressed in the xylem. In a previous study, 41 transcripts were investigated and showed distinct expression profiles in different parts of *S. flavescens* (12). The transcripts related to alkaloid and flavonoid biosynthesis in *S. flavescens* demonstrated organ-specific expression patterns, implying that they might have different physiological processes for biosynthesis, depending on the organ.

The correlation analysis results further showed that 28 and 12 transcripts were positively correlated with the contents of alkaloids and flavonoids, respectively (*R* > 0.8, *P* < 0.05). In the alkaloid biosynthetic pathway, the expression levels of LYS1, AO6, and PMT transcripts were highly and positively correlated with the contents of alkaloids. In a previous study, seven enzyme genes involved in the alkaloid biosynthesis in *S. flavescens* were identified. In the current study, three 4CL, three CHI, two 2′OH, and four CHR were highly and positively correlated with the flavonoids in the flavonoid biosynthetic pathway. PAH was identified as one of the enzymes involved in the flavonoid biosynthesis pathway. These results indicated that these genes resulted in the uneven distribution of alkaloids.

**Materials and methods**

All experimental research and field studies on plants, including the collection of plant material in this study, had complied with relevant institutional, national, and international guidelines and legislation.

**Plant materials.** Three-year-old roots of *S. flavescens* were collected from Wenshan in Yunnan Province at their flowering stage. *S. flavescens* was cultivated with the standard operating procedures established by the Good Agriculture Practices. All roots were carefully washed and separated into three different parts: the periderm, phloem, and xylem (Supplemental file 1: Figure S1). The samples were divided into two parts for metabolite and transcriptome analyses.

**Metabolite analysis.** All of the samples were dried and crushed, and 0.1 g of the powdered sample was weighed and mixed with 1.0 mL of pure methanol under vortex for 1 min and incubated at room temperature for 10 min. The mixture was stored overnight at ~20 °C and centrifuged at 4000 g for 20 min. The upper layer was collected, filtered through a 0.22 μm filter, and transferred to a sample vial. The vial was injected into a column for UPLC-QTOF-MS analysis.

The UPLC-MS analysis was performed using a UPLC system (Waters, UK) coupled to an electrospray ionization-QTOF/MS apparatus (Waters, UK). A 100 mm × 2.1 mm C₁₈ reversed-phase column (Acquity UPLC T3 column, Waters, UK) was used for UPLC separation, and the sample injection volume was 4 μL. The column temperature was kept at 35 °C, and the flow rate was maintained at 0.4 mL/min. The gradient was composed of water containing 0.1% formic acid (A) and acetonitrile containing 0.1% formic acid (B). The linear gradient was set as follows: 0–0.5 min for 5% B, 0.5–7 min for 5%–100% B, 7–8 min for 100% B, and 8–10 min for 5% B. A high-resolution tandem mass spectrometer TripleTOF5600plus (SCIEX, UK) was used to detect metabolites. The Q-TOF was operated in the positive and negative ion modes. The curtain gas was set to 30 PSI, the ion source gas1 was set to 60 PSI, the ion source gas2 was set to 60 PSI, and an interface heater temperature was set at 650 °C. Multivariate data analysis was performed using MetaboAnalyst 4.0 software (http://www.metaboa nalyst.ca/). The PCA was performed to analyze the distribution of samples. One-way ANOVA was used to detect the difference of variance, and variance with FDR ≤ 0.05 was deemed as potential biomarkers. Variable VIP was used to evaluate the variable contribution.

**High-performance liquid chromatography-ultraviolet detection (HPLC-UV) analysis.** The standards of oxymatrine, sophoridine, matrine, trifolirhizin, maackiain, kushenol I, kurarinone, and sophoraflavone G (purity ≥ 98.0%) were purchased from Shanghai Tauto Biotech Company (Shanghai, China). Their batch numbers were 16837–52–8, 6882–68–4, 519–02–8, 6807–83–6, 19908–48–6, 99119–69–4, 34981–26–5, and 97938–30–2, respectively.

The sample extracts were also used for alkaloid and flavonoid quantitative analyses. An Agilent HPLC-UV 1260 series system (Agilent, USA) equipped with a quaternary pump, automatic sampler, column compartment. A VWD was also employed. A 4.6 mm × 250 mm C₁₈ reversed-phase column (with an inner diameter of 5 μm; Eclipse XDB, Agilent, USA) was used for separation, and the sample injection volume was set as 10 μL.
The conditions for alkaloids were set as follows: column temperature of 30 °C, a flow rate of 1.0 mL/min, and a wavelength of 220 nm. The gradient was composed of 80% acetonitrile (A), 10% ethanol (B), and 10% water (C). The conditions for flavonoids were set as follows: column temperature of 35 °C, a flow rate of 1.0 mL/min, and a wavelength of 295 nm. The gradient was composed of acetonitrile (A) and water (B), and the linear gradient was set as follows: 0–25 min for 19%–50% A, 25–30 min for 50%–70% A, 30–40 min for 70% A, and 40–50 min for 70%–40% A.

RNA extraction and illumina sequencing. The total RNA was isolated from different tissues in accordance with the instructions indicated in a plant RNA isolation kit (BioTeke, Beijing, China). The quality of RNA was evaluated on 1% agarose gel, and RNA concentrations were determined with a Nanodrop 2000 spectrophotometer (Thermo Technologies). cDNA library construction and sequencing were performed in accordance with the standards of progress. First, mRNA was enriched from the total RNA by oligo (dT) magnetic beads and broken into short fragments. Then, a random hexamer and RNA fragments were used to prime cDNA synthesis. After purification and connection with adapters, the cDNA library was constructed through PCR amplification. The length of an insert sequence was verified with an Agilent 2100 bioanalyzer system (Agilent Technologies, Santa Clara, CA, USA), and the library was quantified by an ABI Step One Plus real-time PCR system (Applied Biosystems, America). Finally, the qualified cDNA library was sequenced with an Illumina HiSeqTM 2000 system (Illumina Technologies).

Transcriptome analysis. All raw reads were subjected to the cutadapt (v1.9.1) and fqtrim (v0.94) software following quality control to produce clean reads: (1) raw reads including adapter sequences and empty adapter were discarded; (2) reads including unknown N bases comprising more than 5% of the total length were filtered; (3) reads including low-quality bases that comprise more than 20% of the total length were discarded. Then, the indicators of Q20% (sequencing error rate less than 0.01), Q30 (sequencing error rate less than 0.001), and GC% were calculated to evaluate the quality of clean reads. All the 150 bp pair-end RNA-Seq reads were submitted to NCBI (Accession number: PRJNA661972).

De novo assembly was performed in Trinity (v2.4.0) software using 150 bp pair-end reads with default parameters. One assembly was performed using nine sequencing reads, and 58,327 transcripts were obtained. These resultant transcripts were searched against the NCBI nonredundant nucleotide (Nt) database, NCBI nonredundant protein (Nr), and SwissProt protein for functional annotation by using the BLAST algorithm with an E-value cutoff of 1e-35. The functional categories of these unique sequences were further analyzed using the above databases and the KEGG database in BLAST and Blast2GO programs as previously reported in the literature.

The clean reads were mapped to the reference by using Bowtie 2 (v2.2.6) to estimate the expression profiles of the transcripts. The expression levels were calculated with the FPKM by using RSEM software (v1.3.1), and the bowtie parameter was set at mismatch 2. The identification of DEGs was performed using the following criteria: fold change (FC) ≥ 2 and FDR ≤ 0.05. The candidate transcripts involved in the alkaloid and flavonoid biosyntheses were selected in accordance with previous reports and databases with FPKM values of the transcripts converted to log_{20} values (FPKM ≥ 5). They were visualized in a heatmap with pearheatmap package (v1.0.12) in \( R \) to identify the different expression profiles among the three tissues.

Co-expression analysis. The WGCNA was used to analyze the relationships between transcript expressions and component contents with the \( R \) package (v3.2.5). The \( R \) package along with its source code and additional material are freely available at https://horvath.genetics.ucla.edu/html/CoexpressionNetwork/Rpackages/WGCNA/. The network construction and module detection method with default settings were used, including an unsigned topological overlap matrix. All parameters were set as defined: ‘soft_power = 22’, ‘TOM-Type = ‘unsigned’, ‘minModuleSize = 30’, ‘reassignThreshold = 0’, and ‘mergeCutHeight = 0.25’. The \( P \)-value of 0.05 was set as the threshold for a significant correlation.

The candidate transcripts involved in the alkaloid and flavonoid biosyntheses were further selected in accordance with the annotation information to analyze the relationship of transcript expression with alkaloid and flavonoid contents. Pearson correlation coefficient of alkaloid and flavonoid contents with FPKM of transcripts were normalized and then calculated using SPSS (v17.0) software. Pearson correlation bubble chart was constructed with the \( R \) package (v3.5.0) to identify pivotal transcripts related to the contents of alkaloids and flavonoids.

Conclusion
To sum up, the alkaloids and flavonoids showed tissue specificity in \( S. \) flavescens roots. Gene expression profiles also showed tissue specificity. The metabolomes and transcriptomes systematically confirmed the pivotal transcripts regulating the distribution of alkaloids and flavonoids. This study elucidated the mechanism of alkaloids and flavonoids synthesis, accumulation, and transportation, which provide the basis for improving the production of alkaloids and flavonoids through genetic engineering. In addition, these genetic resources could provide comprehensive information on gene discovery, transcriptional regulation, and variety selection for \( S. \) flavescens.

Code availability
All of the transcriptome sequences were submitted to NCBI (Accession number: PRJNA661972).

Received: 14 November 2020; Accepted: 22 March 2021
Published online: 01 April 2021
This study was supported by grants from the Beijing Nova Program (No. Z181100006218020), the Fundamental Research Funds for the Central public welfare research institutes (No. ZZ13-AQ-049 and No. ZKKT17049),
Major Science and Technology Projects of Yunnan Province (No. 2018ZF011) and National Key R&D Plan (No. 2017YFC1702500), National Science and Technology Major Project for “Significant New Drugs Development” (2019ZX09201005-002-001).

**Author contributions**
Conceptualization, D. L. and W. G.; writing-original draft preparation, W. G.; methodology, C. Y.; formal analysis, G. X.; resources, C. Y.; writing-review and editing, D. L.; supervision, C.S., W. J., and D. L.; funding acquisition, C.S. and D. L.; All authors have read and agreed to the published version of the manuscript.

**Competing interests**
The authors declare no competing interests.

**Additional information**

**Supplementary Information** The online version contains supplementary material available at https://doi.org/10.1038/s41598-021-86970-0.

**Correspondence** and requests for materials should be addressed to L.D. or S.C.

**Reprints and permissions information** is available at www.nature.com/reprints.

**Publisher’s note** Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

---

© The Author(s) 2021