Research Article

Kadsura coccinea Lignan Metabolism Based on Metabolome and Transcriptome Analysis

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Kadsura coccinea is an important resource of traditional Chinese medicine. We find out the gene information of enzymes related to lignan biosynthesis and metabolism of Kadsura coccinea, so as to provide a scientific basis for the breeding of new varieties of Kadsura coccinea. In this paper, 2-year-old Kadsura coccinea from Hunan Kadsura coccinea provincial germplasm resource bank was used as the material and its root, stem, and leaf were analyzed by extensive targeted metabolomics combined with transcriptome sequencing. The results showed the following: (1) 51 lignans were detected by metabolome analysis, and the content of lignans in roots was higher than that in stems and leaves. The high content of lignans in roots, stems, and leaves includes ring-opening isolarch phenol-4-o-glucoside, narrow leaf schisandrin E, and schisandrin B. (2) After transcriptome sequencing, 13 classes of 137 Unigenes related to lignan biosynthesis pathway were retrieved. The analysis of differential genes in different parts showed that the overall expression amount and species of Kadsura coccinea lignan synthase gene in stems and leaves were closer than those in roots. CCoAOMT, C3H, and SIDR gene families are mainly expressed in roots and stems. (3) Metabolome combined with transcriptome analysis further screened these genes and obtained 11 genes of enzyme gene families such as HCT, DIR, COMT, CAD, SIDR, and PLR, which are highly correlated in lignan synthesis. Therefore, there are many lignans and their synthase-related genes in Kadsura coccinea roots, stems, and leaves, but the content and expression of different lignans and their synthase-related genes are quite different in each part. In this study, the gene information of the Kadsura coccinea lignan biosynthesis enzyme was obtained for the first time, which laid a good foundation for the cloning and molecular breeding of the key enzyme gene of lignan biosynthesis.

1. Introduction

Kadsura coccinea is an evergreen woody vine of the Schisandra genus in the Schisandra family, also known as cold rice ball, bufuna, chicken intestine wind, etc. Kadsura coccinea is an important medicinal resource, mainly distributed in Hunan, Guangxi, Guizhou, Fujian, and other places [1]. Folk medicine mostly uses its roots, vines, and fruits to treat stomach diseases, rheumatoid arthritis, bruises, swelling and pain, irregular menstruation, and so on [2]. Pharmacological studies showed that Kadsura coccinea has the effects of antitumor [1], antioxidation [3], liver protection [4], and inhibition of acetylcholinesterase [5].

The main active components of Kadsura coccinea are lignans, triterpenes, sesquiterpenes, sterols, and other compounds, among which lignans are the main active components [6]. At present, 121 kinds of lignans have been isolated and identified, mainly including biphenyl cyclooctadiene type, spirobenzofuran type, and 6,9-oxbiphenyl cyclooctadiene type, followed by a small amount of arylaminophthalene type and dibenzylbutane type lignans [7]. Guo Xia et al. [8] measured the highest content of total lignans in Huaihua, Hunan Province, from Kadsura coccinea varieties collected from six different provenances. At present, the research on Kadsura coccinea mainly focuses on chemical composition analysis, pharmacological activity
screening, and cultivation [9]. However, the analysis of the *Kadsura coccinea* secondary metabolic pathway has not been carried out, and the discovery of key enzyme genes is also rare. With the development of gene sequencing, based on extensive targeted metabolome combined with transcriptome sequencing analysis, studying the biosynthetic regulation pathway of *Kadsura coccinea* lignans will contribute to the development of molecular markers of *Kadsura coccinea*, carry out molecular breeding, cultivate excellent *Kadsura coccinea* varieties with high lignan content, and lay a foundation for the further development and utilization of *Kadsura coccinea* traditional Chinese medicine resources.

2. Materials and Methods

2.1. Materials. The experimental material *Kadsura coccinea* comes from the provincial Germplasm Resource Bank of *Kadsura coccinea* in Hunan Province. Six duplicate tissue materials were collected from the root, stem, and leaf of *Kadsura coccinea* introduced from Huaihua, Hunan Province. After washing and drying with sterilized deionized water, they were put into a 2 mL cryopreservation tube. After quick freezing with liquid nitrogen, they were stored in a low-temperature refrigerator at −80°C for standby.

2.2. Method

2.2.1. Metabolomic Analysis. Three biological replicates were taken from the root, stem, and leaf of *Kadsura coccinea* preserved at ultra-low temperature, and a total of 9 sample materials were analyzed for extensive targeted metabolome.

(1) Sample Metabolite Extraction. Nine samples of the root, stem, and leaf were taken, and the three of each were placed in a freeze dryer (Scientz-100F) for vacuum freeze-drying for 30 min, repeat the process 6 times, and place 70% methanol extract. We vortex the extract for 30 s, hold when, we weigh 100 mg of powder and dissolve it in 1.2 mL of ultrapure water, they were put into a 2 mL cryopreservation tube. After quickly freezing with liquid nitrogen, they were stored in a low-temperature refrigerator at −80°C for standby.

(2) UPLC-MS/MS Analysis.

- **Liquid phase conditions:**
  - Chromatographic column: Agilent SB-C18 1.8 μm, 2.1 mm * 100 mm
  - Mobile phase: phase A is ultrapure water (add 0.1% formic acid), and phase B is acetonitrile (add 0.1% formic acid)
  - Elution gradient: the proportion of phase B is 5% in 0.00 min, the proportion of phase B increases linearly to 95% in 9.00 min and remains at 95% for 1 min, for 10.00–11.10 min, and the proportion of phase B decreases to 5% and balances with 5% to 14 min
  - Flow rate: 0.35 mL/min; column temperature: 40°C; injection volume: 4 μL

(2) Mass spectrum conditions:

Linear ion trap and three-pole quadrupole scanning were performed on a three-pole quadrupole linear ion trap mass spectrometer (Q TRAP), AB4500 Q TRAP UPLC/MS/MS system. The system is equipped with an electrospray ionization source ion spray interface, which can be controlled using Analyst 1.6.3 software (AB Scie) to obtain two positive and negative ion modes. The operating parameters of the electrospray ion source are as follows: ion source, turbine spray; source temperature of 550°C; ion spray voltage of 5500 V (positive ion mode)/–4500 V (negative ion mode). The ion source gas I, ion source gas II, and curtain gas are set to 50, 60, and 25.0 psi, respectively, and the collision-induced ionization parameter is set to high value. In triple quadrupole and linear ion trap modes, 10 and 100 μmol/L are used, respectively. The instrument tuning and mass calibration were carried out with mol/L polypropylene glycol solution. Triple quadrupole scanning was performed using multireaction monitoring mode and by setting the collision gas (nitrogen) to medium. By further optimizing the declustering voltage and collision energy, the declustering voltage and collision energy of each multireaction monitoring ion pair are obtained. According to the metabolites eluted in each period, a specific set of multireaction monitoring ion pairs are monitored in each period [9].

2.2.2. Screening of Lignans-Related Components. After the samples from different parts of *Kadsura coccinea* were analyzed by UPLC-MS/MS, based on the self-built database of Baimike Technology Co. Ltd., the related components of lignans in *Kadsura coccinea* root, stem, and leaf were analyzed qualitatively and quantitatively according to the secondary spectrum information.

2.2.3. Transcriptome Sequencing Analysis

(1) Construction of Sequencing Library and Computer Sequencing. The trizol method (Tiangen dp411 Kit) was used to extract the total RNA of 9 samples from the root, stem, and leaf of *Kadsura coccinea* for 3 biological repetitions. After passing the nucleic acid concentration and integrity test, the second-generation transcriptome sequencing library was constructed by using VAHTS Universal V6 RNA-seq Library Prep Kit for Illumina. It includes magnetic bead enrichment, mRNA reverse transcription to form cDNA, sequencing connector connection, PCR amplification, and enrichment of samples. After the library passed the quality inspection, the transcriptome of 9 samples was sequenced through Illumina NovaSeq 6000 platform.
Components.

3. Results and Analysis

3.1. Metabonomic Analysis of Kadsura coccinea-Related Components. We take Kadsura coccinea root, stem, and leaf for 3 biological replicates, respectively, and a total of 9 sample materials for metabolomic analysis. Based on the UPLC-MS/MS detection platform and the database established by Baimaike Technology Co., Ltd., a total of 1000 metabolites were detected. These metabolites are divided into 13 categories, including 172 phenolic acids, 166 lipids, 134 flavonoids, 96 organic acids, 86 amino acids and their derivatives, 66 terpenes, 60 nucleotides and their derivatives, and 55 lignans and coumarins, as shown in Table 1. The contents of lignans, terpenoids, and alkaloids in roots were higher than those in stems and leaves. The content of flavonoids in roots was lower than that in stems and leaves.

3.2. Analysis of Lignans-Related Components. Among the 55 primary metabolites of lignans and coumarins detected, there were 51 secondary metabolites of lignans. The lignans with high content in Kadsura coccinea roots are ring-opening isolarch phenol-4-O-glucoside, isoschisandrin B, mangliesin D, etc., as shown in Figure 1.

Lignans with high content in Kadsura coccinea stem include schisandrin E, ring-opening isolarch-9′-O-glucoside, schisandrin ethyl, etc., as shown in Figure 2.

The high content of lignans in Kadsura coccinea leaves includes schisandrin B, ring-opening isolarch-9′-O-glucoside, inulin C, etc., as shown in Figure 3.

3.3. Transcriptome Sequencing Results. Transcriptome sequencing was performed on 9 samples from Kadsura coccinea root, stem, and leaf. A total of 60.44 Gb Clean Data was obtained. The Clean Data of each sample reached 5.73 Gb, and the percentage of Q30 base was 94.35% or more. After assembly, a total of 54309 Unigenes were obtained. Among them, there are 16141 Unigenes with a length of more than 1 kb. The functional annotation of Unigenes includes the comparison with cog, go, KEGG, KOG, Pfam, Swiss-Prot, TrEMBL, eggNOG, and NR databases, as shown in Table 2. A total of 26658 Unigene annotation results are obtained, of which 13683 Unigenes with a length of more than 1000 bases are annotated. Among all databases, the number of Unigenes annotated in the nonredundant protein (NR) database is the largest, reaching 25852, 12302 for 300–1000 and 13550 for more than 1000, as shown in Table 2.

3.4. Gene Analysis of Enzymes Involved in Lignans Synthesis. Combined with the research on lignan biosynthesis of other species, we continued to excavate the enzyme genes related to lignan biosynthesis from the 26658 Unigenes annotated. Referring to the phenylpropane metabolic pathway (ko00940) and some other reported lignan biosynthesis pathways (ko00998) [10–15], a total of 13 classes of 137

Table 1: The average value of species, quantity, and relative content of metabolites in roots, stems, and leaves of Kadsura coccinea.

<table>
<thead>
<tr>
<th>Primary metabolites</th>
<th>Number</th>
<th>Root</th>
<th>Stem</th>
<th>Leaf</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lignans and coumarins</td>
<td>55</td>
<td>7.57E-02</td>
<td>4.85E-02</td>
<td>4.74E-02</td>
</tr>
<tr>
<td>Terpenoids</td>
<td>66</td>
<td>4.95E-02</td>
<td>2.80E-02</td>
<td>3.74E-02</td>
</tr>
<tr>
<td>Flavonoids</td>
<td>134</td>
<td>5.18E-02</td>
<td>1.32E-01</td>
<td>2.47E-01</td>
</tr>
<tr>
<td>Phenolic acids</td>
<td>172</td>
<td>1.56E-01</td>
<td>2.14E-01</td>
<td>2.21E-01</td>
</tr>
<tr>
<td>Lipid</td>
<td>166</td>
<td>1.23E-01</td>
<td>1.22E-01</td>
<td>1.24E-01</td>
</tr>
<tr>
<td>Organic acid</td>
<td>134</td>
<td>5.31E-02</td>
<td>7.11E-02</td>
<td>4.63E-02</td>
</tr>
<tr>
<td>Amino acids and their derivatives</td>
<td>86</td>
<td>2.17E-01</td>
<td>1.63E-01</td>
<td>8.52E-02</td>
</tr>
<tr>
<td>Nucleotides and their derivatives</td>
<td>60</td>
<td>1.41E-02</td>
<td>1.82E-02</td>
<td>2.20E-02</td>
</tr>
<tr>
<td>Alkaloid</td>
<td>26</td>
<td>1.35E-01</td>
<td>5.66E-02</td>
<td>2.78E-02</td>
</tr>
<tr>
<td>Tannin</td>
<td>12</td>
<td>2.61E-02</td>
<td>1.30E-02</td>
<td>3.84E-03</td>
</tr>
<tr>
<td>Other classes</td>
<td>127</td>
<td>9.85E-02</td>
<td>1.34E-01</td>
<td>1.38E-01</td>
</tr>
</tbody>
</table>

(2) Bioinformatics Analysis of Transcriptome Sequencing. We filter the original data obtained, remove the joint sequence and low-quality reading sequence, and obtain high-quality reading sequence data. The data were assembled by Trinity software to obtain the UniGene Library of Kadsura coccinea root, stem, and leaf tissues. The quality of the sequencing library was evaluated by examining the randomness of mRNA fragmentation, the dispersion of the length of inserted fragments, the library capacity, and the adequacy of mapped reads compared to the UniGene library. The qualified database then carries out functional annotation and expression analysis on UniGene through BLAST and other software.
Unigenes related to the lignan biosynthesis pathway were retrieved, of which the gene family with a large number is hydroxycinnamoyl transferase (HCT) polymerization protease (DIR) and cinnamoyl CoA reductase (CCR), 37, 21, and 16. Cinnamic acid-4-hydroxylase (C4H), coumaric acid-3-hydroxylase (C3H), and phenylalanine ammonia-lyase (PAL), respectively, have a small number of gene families, as shown in Table 3.

Furthermore, we analyze the relative gene expression of the 137 Unigenes related to lignan synthesis in 9
transcriptome samples of *Kadsura coccinea* root, stem, and leaf (3 replicates in each part) and cluster-analyze the differential genes, as shown in Figure 4. Heat map analysis showed that the total expression and species of *Kadsura coccinea* lignan synthase gene in stems and leaves were closer to those in roots. Caffeoyl COA oxi-methyltransferase (CCoAOMT), coumaric acid-3-hydroxylase (C3H), and isolarch dehydrogenase (SIDR) are among several enzyme gene families, which are mainly expressed in roots and stems.

### 3.5. “Transcription + Metabolism” Correlation Analysis of Lignan Biosynthesis Pathway.

Among the 137 Unigenes related to lignan synthesis obtained from the transcriptome, 30 Unigenes with high expression were screened. The correlation analysis was carried out in the phenylpropane metabolic pathway (ko00940) with the three lignans with the highest content in roots, stems, and leaves obtained from the metabolome analysis.

In the process of root and stem comparison, 13 differential gene expressions were related to ring-opening isolarch...
phenol-4-o-glucoside and schisandrin E. Among them, there are 8 positive correlations and 4 negative correlations involving ring-opening isolarch phenol-4-o-glucoside. Among the positive correlations, hct9, dir15, comt4, and cad3 have relatively large correlations, as shown in Figure 5.

During the comparison of root and leaf, 13 differential gene expressions were related to schisandrin B, including 4 positive and 9 negative correlations. The four positive correlation genes were HCT9, HCT10, CAD3, and HCT1, as shown in Figure 6.

In the comparison of stem and leaf, 17 differential gene expressions were related to schisandrin E and schisandrin B. Schisandrin E has 11 positive correlations and 6 negative correlations. In the positive correlations, COMT4, DIR15, PLR4, and SIDR2 are more correlated; Schisandrin B has 6 positive correlations and 11 negative correlations. Among the positive correlations, HCT10, DIR4, and CAD3 genes have relatively large correlations, as shown in Figure 7.

### 4. Discussion

Lignans are important secondary metabolites of plants, which play an important role in insect resistance and stress growth of plants. At the same time, lignans also have important pharmacological activities such as antitumor, anti-HIV, anti-inflammatory, liver protection, and antioxidation. The research on chemical constituents of *Kadsura coccinea* has made rapid progress, but it mainly focuses on chemical composition analysis and pharmacological activity research. At present, the research on the lignan biosynthesis pathway of *Kadsura coccinea* has not been reported. In the research field of lignan biosynthesis in *Kadsura coccinea*, our research team used the method of metabolome combined with transcriptome analysis for the first time.

At present, the metabolomic analysis of *Kadsura coccinea* focuses on the study of relevant components of fruits and seeds or the study of flavonoids in different parts. However, the metabolomic study of lignans, the main pharmacodynamic component of *Kadsura coccinea*, has not been carried out.

However, lignans and their derivatives are one of the main chemical constituents isolated from black tigers. Lu et al. [18] used UV-vis spectrophotometry to determine the content of total lignan in black tigers from three different producing areas in Guangxi Province and found that the content of total lignan in black tigers from different producing areas was different, with the content between 0.93% and 2.11%. The main lignan compounds isolated from *Schisandra chinensis* are biphenyl cyclooctene lignans. At present, there are 49 biphenyl cyclooctadiene lignans isolated and identified from black tiger, which can be divided into three categories: the common biphenyl cyclooctadiene type, spirobenzofurane biphenyl cyclooctadiene type, and 6,9 oxbridge biphenyl cyclooctadiene type lignans. Through ultra-high-performance liquid chromatography-tandem mass spectrometry, the research team analyzed the related components of lignans in the root, stem, and leaf of *Kadsura coccinea* and obtained a total of 51 lignans, and the content in the root is higher than that in the stem and leaf. The high content of lignans in roots, stems, and leaves includes ring-opening isolarch phenol-4-o-glucoside, narrow leaf schisandrin E, and schisandrin B. This study enriches the

<table>
<thead>
<tr>
<th>Gene family</th>
<th>Number of Unigenes</th>
<th>Code ID</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cinnamyl alcohol dehydrogenase (CAD)</td>
<td>11</td>
<td>c138243, c127253, c100870, c100870, c132969, c134622, c75605, c139976, c142940, c106854, c139288</td>
</tr>
<tr>
<td>Cinnamate-4-hydroxylase (C4H)</td>
<td>3</td>
<td>c133158, c141289, c140162</td>
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<tr>
<td>Caffeoyl COA oxymethyltransferase (CCoAOMT)</td>
<td>5</td>
<td>c123906, c147567, c130267, c134322, c139948</td>
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<tr>
<td>4-Coumarinyl CoA ligase (4CL)</td>
<td>11</td>
<td>c141218, c148238, c148167, c139095, c105572, c147311, c152599, c147707, c148297, c141218, c142717</td>
</tr>
<tr>
<td>Cinnamoyl CoA reductase (CCR)</td>
<td>16</td>
<td>c141261, c135703, c120215, c140397, c14246, c146949, c136448, c140375, c138246, c141087, c134396, c147312, c144833, c149012, c140534, c129111</td>
</tr>
<tr>
<td>Coumaric acid-3-hydroxylase (C3H)</td>
<td>3</td>
<td>c138815, c130179, c145761</td>
</tr>
<tr>
<td>Phenylalanine ammonia-lyase (PAL)</td>
<td>3</td>
<td>c145137, c142300, c125745</td>
</tr>
<tr>
<td>Hydroxycinnamoyl transferase (HCT)</td>
<td>37</td>
<td>c148893, c149025, c143322, c137863, c135415, c132343, c138259, c137078, c130160, c135415, c145853, c132343, c150077, c139674, c136799, c149948, c12859, c134687, c133462, c146001, c136937, c144136, c136395, c146817, c149992, c140186, c142265, c145173, c131394, c136937, c93393, c148160, c125352, c135415, c132085, c135897, c146817</td>
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<tr>
<td>Caffeic acid oxymethyltransferase (COMT)</td>
<td>6</td>
<td>c146701, c106776, c122011, c144220, c149283, c125532, c135415, c132085, c135897, c146817</td>
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<tr>
<td>Isolaricin dehydrogenase (SIDR)</td>
<td>10</td>
<td>c127415, c145168, c114390, c145090, c135982, c128786, c130422, c132278, c142699, c126873</td>
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<tr>
<td>Polymerized protease</td>
<td>21</td>
<td>c138256, c182448, c122470, c135239, c131102, c134134, c134002, c114144, c134644, c135894, c121895, c126128, c132052, c135894, c100228, c125953, c112255, c133684, c144151</td>
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<td>Terpineol reductase</td>
<td>4</td>
<td>c138332, c140550, c145666, c147861</td>
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<tr>
<td>Caffeoyl shikimate esterase (CSE)</td>
<td>7</td>
<td>c139639, c148145, c142263, c133532, c141052, c139096, c134673</td>
</tr>
</tbody>
</table>

Table 3: The major lignan synthase genes in *Kadsura coccinea*. 

### Table 3: The major lignan synthase genes in *Kadsura coccinea*.
Figure 4: The relative expression of lignan synthase gene in *Kadsura coccinea* roots, stems, and leaves.

Figure 5: Correlation network of differentially expressed genes and metabolites in *Kadsura coccinea* lignan biosynthesis (root and stem).

Figure 6: Correlation network of differentially expressed genes and metabolites in *Kadsura coccinea* lignan biosynthesis (root and leaf).
study of lignans in different parts of *Kadsura coccinea* and lays a solid foundation for the further development of lignans in different parts of *Kadsura coccinea*.

At present, there is no report on the study of enzyme genes related to lignan biosynthesis in *Kadsura coccinea*. After obtaining UniGene data information through transcriptome sequencing, according to the reported enzymes involved in lignan biosynthesis and metabolism in other plants, this study further searched the information related to lignans synthesis in *Kadsura coccinea* and a total of 13 classes of 137 Unigenes related to lignan biosynthesis pathway were retrieved. Among them, CAD, C4H, CCoAOMT, 4CL, CCR, C3H, PAL, HCT, COMT, and CSE belong to the upstream stage of lignan biosynthesis, which is responsible for the production of lignan precursor coniferol from phenylalanine through a series of reactions. DIR, PLR, and SIDR are lignans such as ring-opening isolarch alcohol and Moha-none, which are further generated from coniferol through turpentine alcohol and larch alcohol. The analysis of differential genes in different parts showed that the overall expression and species of *Kadsura coccinea* lignan synthase gene in stems and leaves were closer than those in roots. CCoAOMT, C3H, and SIDR gene families are mainly expressed in roots and stems. Furthermore, we analyze the transcriptional groups of 11 genes related to HCRD and PLT. In this study, the gene information of the *Kadsura coccinea* lignan biosynthesis enzyme was obtained for the first time, which laid a good foundation for the cloning and molecular breeding of key enzyme genes of *Kadsura coccinea* lignan biosynthesis.

5. Conclusion

The *Kadsura coccinea* has great economic value, and it has become a characteristic industrial source in Hunan, Fujian, and other provinces. However, due to the lack of research and development on black tiger, there are not many kinds of black tiger-related products with low added value and industrial development is restricted to a certain extent. Although we have a comprehensive understanding of the chemical composition of the black tiger, the study of its efficacy and the basis of its pharmacological substances is still very weak. Firstly, many chemical components still lack activity and mechanism research. Secondly, pharmacological studies are not comprehensive, in-depth, and systematic, which cannot provide sufficient support for the clinical application and efficacy development of the black tiger. Thirdly, the quantitative study of its chemical composition is obviously lacking, which leads to the great limitation of the standard study. Finally, there is a lack of metabolism and
metabolite activity in in vivo research. These problems need to be focused on and solved in future research work.

Data Availability
The data used to support the findings of this study are available from the corresponding author upon request.

Conflicts of Interest
The authors declare that they have no conflicts of interest.

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